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71) Applicant: EUROPEAN ECONOMIC COMMUNITY E.E.C.
Bâtiment Jean Monnet
Plateau du Kirchberg
L-2920 Luxembourg (LU)

(2) Inventor: Klein, Jürgen Robert Mackenbacher Strasse 15 D-67685 Weilerbach (DE) Inventor: Plapp, Roland Am Harzhübel 88 D-67663 Kaiserslautern (DE)

(74) Representative: Grosset-Fournier, Chantal Catherine
Grosset-Fournier & Demachy s.a.r.l.
103 rue La Fayette
F-75010 Paris (FR)

(54) Lys-aminopeptidase PepN from lactobacillus delbruckii ssp. Lactis, nucleic acids coding for it, and its use in fermentation processes.

57 The invention relates to a protein as produced by *Lactobacillus delbrūckii* ssp. *lactis*, and capable of hydrolysing β-naphtylamides, or fragments thereof having this enzymatic activity.

The invention relates also to the use of said protein or fragments thereof for the preparation of fermented foodstuff, and more particularly of cheese.

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The invention relates to a protein, PepN, as produced by Lactobacillus delbrūckii ssp. lactis and capable of hydrolysing β -naphtylamides, or polypeptides derived thereof, and more particularly recombinant PepN from said Lactobacillus strain or recombinant polypeptides derived thereof, said derived polypeptides having this enzymatic activity of hydrolysing β -naphtylamides.

The invention relates also to the use of said protein or polypeptides derived thereof in fermentation processes, such as processes for the preparation of fermented foodstuff, and more particularly cheese.

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The invention also relates to processes for preparing the PepN from said *Lactobacillus* strain or said derived polypeptides, which are in a state of biological purity such that they can be used in fermentation processes.

The invention also relates to nucleic acids (or nucleotide sequences) coding for PepN from said *Lactoba-cillus* strain or said derived polypeptides, and to cellular hosts containing said nucleic acids and their use in fermentation processes.

Furthermore, the invention relates to fermentation processes, and more particularly to processes for the preparation of fermented foodstuff, and kits, using the PepN from said *Lactobacillus* strain or said derived polypeptides, and/or said cellular hosts.

By "recombinant polypeptides" it is to be understood that it relates to any molecule having a polypeptidic chain liable to be produced by genetic engineering, through transcription and translation of a corresponding DNA sequence under the control of appropriate regulation elements within a efficient cellular host. Consequently the expression "recombinant polypeptides" such as is used herein does not exclude the possibility for the polypeptides to comprise other groups, such as glycosylated groups.

The term "recombinant" indeed involves the fact that the polypeptide has been produced by genetic engineering, particularly because it results from the expression in a cellular host of the corresponding nucleic acid sequences which have previously been introduced into the expression vector used in said host.

Nevertheless, it must be understood that this expression does not exclude the possibility for the polypeptide to be produced by a different process, for instance by classical chemical synthesis according to methods used in the protein synthesis or by proteolytic cleavage of larger molecules.

The expression "biologically pure" or "biological purity" or "in a substantially pure form" means on the one hand a grade of purity such that the polypeptide or recombinant polypeptide can be used for the preparation of fermented foodstuff and on the other hand the absence of contaminants, more particularly of natural contaminants.

The lactobacilli used as starter cultures in the industrial dairy fermentation need to have an efficient proteolytic system. Since the concentration of free essential amino acids in milk are low, a variety of proteolytic enzymes is necessary for the breakdown of milk casein. By the combined action of proteinases and peptidases, milk protein is degraded to peptides and amino acids, which are required for cell growth and which contribute to the organoleptic properties of the foods. Also the flavour of milk products is primarily based on this degradation. The proteolytic cascade must be initiated extracellularly and successive coordinated degradation of peptides by endo- and exopeptidases, with cell wall and cytoplasmic localizations, generate amino acids necessary for growth. The importance of the proteolytic system for dairy product quality has resulted in an increased fundamental research of the enzymes and genes involved. Various genes and corresponding enzymes from lactococci strains have been investigated, but less is known about the proteolytic system of lactobacilli.

An aspect of the invention is to provide a new family of nucleic acids coding for new proteins and polypeptides which can be used in fermentation processes, and more particularly for the preparation of fermented foodstuff

Another aspect of the invention is to provide cellular hosts transformed with said nucleic acids which can be used in fermentation processes, and more particularly as starter organisms in the fermentation of milk.

Another aspect of the invention is to provide nucleic acids coding for the peptidic chain of biologically pure recombinant polypeptides which enable their preparation on a large scale.

Another aspect of the invention is to provide new proteins and polypeptides which can be used in fermentation processes.

Another aspect of the invention is to provide new fermentation processes, such as processes for the preparation of fermented foodstuff, and more particularly cheese.

The invention relates to the cloning, expression, and nucleotide sequence of the lys-aminopeptidase gene (pepN) from Lb. $delbr\bar{u}ckii$ ssp. lactis and to some characteristics of the purified enzyme. The gene has been designated pepN since it complements an E. $coli\ pepN$ mutation, and as reported hereafter, there is extensive amino acid homology to other aminopeptidases N (E.C. 3.4.11.2). The enzyme PepN is characterized by its specific action on β -naphtylamides.

The invention relates to a protein in a substantially pure form, as produced by *Lactobacillus delbrūckii* ssp. *lactis*, said protein, also called PepN, being capable of hydrolysing β-naphtylamides, and more particularly ly-

sine- β -naphtylamide, or fragments thereof having this enzymatic activity (i.e. this hydrolysing amino acid β -naphtylamides activity).

By hydrolysing amino acid β -naphtylamides, one should understand that such activity corresponds to aminopeptidase N (PepN); see Table 3 hereafter.

The invention relates more particularly to the PepN protein such as described above and produced by *Lactobacillus delbrückii* ssp. *lactis* WS87, deposited at the Deutsche Sammlung von Mikroorganismen (DSM) under the number 7290 on October 15, 1992.

The invention relates more particularly to the PepN protein represented on figure 3, and by SEQ ID NO 2, or polypeptides derived thereof, such as fragments or muteins (which differ from said protein by addition and/or substitution and/or suppression of one or several amino acid) thereof, provided that said derived polypeptides are capable of hydrolysing β -naphtylamides.

Said PepN represented by SEQ ID NO 2, is more particularly characterized in that:

- its isoelectric point calculated from the nucleotide sequence represented by SEQ ID NO 1, is 4,48, and its isoelectric point determined by preparative isoelectric focusing after purification of said protein, is 4.2.
- its molecular weight calculated from the nucleotide sequence represented by SEQ ID NO 1, is 95,358 kDa, and its molecular weight determined after purification, is 95 kDa,
- its specific chromogenic substrates are chosen among those listed in Table 3,
- its specific inhibitors are the following:
 - . EDTA: end concentration 1 mM: 3 % relative activity,
 - . Phenanthroline: end concentration 1 mM: 4 % relative activity.

The percentages of relative activity indicated above, correspond to residual activities after addition of the inhibitor as compared with the control without inhibitor. The conditions and methods for determination are taken from "Proteolytic enzymes, a practical approach, Beynon R.J. and Bond J.S., Oxford University Press 1989." (Chapter 4, Determination of protease mechanism, Ben M. Dunn).

The PepN according to the invention is more particularly characterized in that it is obtained in a purified state, from a cell extract of said *Lactobacillus delbrückii* ssp. *lactis* WS87 by the following procedure:

- fractionation of the cell extract by salting out at 4°C with streptomycin sulfate,
- centrifugation,
- anion exchange chromatography by applying the supernatant obtained at the previous step to an appropriate column, such as a column of Q-Sepharose Fast Flow (Pharmacia), and pooling the eluted fractions having the highest specific activity against L-Pro-p-nitroanilide,
- preparative isoelectric focusing, by applying the pooled fractions obtained at the previous step to an appropriate column, such as a LKB column (Types 8100-1), with ampholytes within a range of pH 4 to 6, and pooling the eluted fractions having the highest specific activity against Lys-paranitroanilide.

The invention also relates to nucleic acids coding for a protein according to the invention, i.e. for the PepN protein or plypeptides derived thereof as described above.

The invention relates more particularly to nucleic acids characterized in that:

- they comprise all or part of the nucleic acid represented on figure 3, and by SEQ ID NO 1, coding for a protein according to the invention, or its complementary sequence,
- or they hybridize with all or part of said nucleic acid represented by SEQ ID NO 1, or with its complementary sequence.

The invention relates more particularly to a nucleic acid coding for the PepN protein represented by SEQ ID NO 2, or for polypeptides derived thereof having said hydrolysing β -naphtylamides activity, said nucleic acid comprising all or part of the nucleotide sequence delimited by the nucleotide located in position 316 and the nucleotide located in position 2844 in the nucleotide sequence represented by SEQ ID NO 1.

The invention also relates to any nucleic acid susceptible to hybridize with all or part of nucleic acids such as described above, and more particularly to any nucleic acid derived from this latter according to the degeneracy of the genetic code, and more particularly by substitution, and/or addition, and/or suppression of one or several nucleotides of said nucleic acids described above, said derived nucleic acid being able to code for the PepN protein represented by SEQ ID NO 2, or for polypeptides derived thereof such as defined above.

By way of illustration, the hybridization above-mentioned can be performed with membranes like Heybond™-N (Amersham) or Hybridization Transfer Membrane (Micron Separations Inc.) under conditions as described by the supplier.

The invention also relates to recombinant nucleic acid containing at least one of the nucleic acids according to the invention, inserted in a heterologous nucleic acid.

The invention also relates to recombinant vectors comprising a vector sequence, notably of the type plasmid (such as plasmids originating from lactic acid bacteria), cosmid or phage, and a nucleic acid described

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above, in one of the non-essential sites for its replication, and optionally one or several nucleic acid(s) coding for other aminopeptidases such as PepX (Meyer-Barton et al., 1993).

Prefered recombinant vector contains in one of its non essential sites for its replication necessary elements to promote the expression of polypeptides according to the invention, in a cellular host and possibly a promoter recognized by the polymerase of the cellular host, particularly an inducible promoter and possibly a signal sequence and/or an anchoring sequence.

The invention relates more particularly to recombinant vector, such as described above, containing the elements enabling the expression by *E. coli* or lactic acid bacteria including *lactococcus* and, especially, lactobacilli of the thermophilic group, of a nucleic acid according to the invention, inserted in the vectors.

The invention relates more particularly to recombinant vector, such as described above, constructed from replicons isolated from lactobacilli, especially from thermophilic *Lactobacillus* species.

The invention also relates to cellular hosts which are transformed by a recombinant vector according to the invention, and comprising the regulation elements enabling the expression of the nucleotide sequence coding for a protein according to the invention, and optionally, other aminopeptidases such as PepX, in these hosts.

The invention relates more particularly to cellular hosts, such as described above, chosen from among bacteria such as *E.coli*, or lactic acid bacteria including *lactococcus* and, especially, lactobacilli of the thermophilic group, transformed by a vector according to the invention.

From the nucleic acids of the invention, probes (i.e. cloned or synthetic oligonucleotides) can be inferred. These probes can be from 25 nucleotides up to the total length of the *pepN* gene. These oligonucleotides can also be used as amplification primers in the PCR technique (PCR, Mullis and Faloona, Methods in Enzymology, vol. 155, p. 335, 1987) to generate specific enzymatically amplified fragments and/or as probes to detect fragments amplified between bracketing oligonucleotide primers.

The invention also relates to the expression products of a nucleic acid expressed by a transformed cellular host according to the invention.

The invention also relates to a process for preparing the PepN protein such as described above, comprising the following steps:

- the culture in an appropriate medium of a cellular host according to the invention,

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- the recovery of the polypeptide produced by the above said cellular host from the above said culture medium,
- the purification of the protein thus obtained, more particularly according to the method described above.
 The invention also relates to fermentation processes comprising a step of treatment of material to be fermented with:
 - an appropriate amount of the PepN protein or polypeptides derived thereof according to the invention, and optionally other proteases, and more particularly aminopeptidases such as PepX, and/or
 - an appropriate amount of at least one of the transformed cellular host such as described above, comprising a nucleotide sequence coding for PepN, and optionally for other proteases, and more particularly aminopeptidases such as PepX,
 - and optionally, an appropriate amount of lactobacilli, such as Lactobacillus delbrückii ssp. lactis.

The invention also relates to a process for the preparation of fermented foodstuff, and more particularly of cheese, which comprises a step of treatment of food material to be fermented, such as milk, with:

- an appropriate amount of the PepN protein polypeptides derived thereof according to the invention, and optionally other proteases, and more particularly aminopeptidases such as PepX,
- an appropriate amount of at least one of the transformed cellular host such as described above, comprising a nucleotide sequence coding for PepN, and optionally for other proteases, and more particularly aminopeptidases such as PepX,
- and optionally, an appropriate amount of lactobacilli, such as Lactobacillus delbrückii ssp. lactis.

The process for the preparation of fermented foodstuff described above can also comprise a step of treatment of food material to be fermented, such as milk, with other species and strains susceptible to be used as starter organisms in fermentation processes, and more particularly lactic acid bacteria susceptible to produce a PepX protein, such as Lactococcus, Streptococcus and Lactobacillus.

The fermentation processes according to the invention, are more particularly characterized in that they can be used as fermentation processes for the obtention of hard cheeses, such as Emmentaler type cheese (South Germany, Switzerland).

The invention also relates to foodstuff, and more particularly cheeses, such as obtained by fermentation processes as described above.

The invention also relates to the use of all or part of nucleotide sequences described above, as tools for analytical purpose. In that respect, the invention relates more particularly to the use of the above-mentioned nucleotide sequences, and more particularly all or part of the SEQ ID NO 1 sequence, in procedures for amino

acid sequencing of peptides and production of chemical or pharmaceutical compounds in which the hydrolysis of β -naphtylamide residues is a required step.

The invention relates more particularly to the use of the polypeptides described above for the determination of the exact localization of PepN in *Lactobacillus delbrückii* ssp. *lactis*, rendering possible experiments performed by immunoblotting after cell fractionation and by electron microscopy of immunogold labelled peptidases.

The polypeptides of the invention can also be prepared according to the classical techniques in the field of peptide synthesis.

The synthesis can be carried out in homogeneous solution or in solid phase.

For instance, the synthesis technique in homogeneous solution which can be used is the one described by Houbenweyl in the book entitled "Methoden der organischen Chemie" (Method of organic chemistry) edited by E. Wünsch, vol. 15-I and II, THIEME, Stuttgart 1974.

The polypeptides of the invention can also be prepared in solid phase according to the methods described by Atherton and Sheppard in their book entitled "Solid phase peptide synthesis" (IRL Press, Oxford, New York, Tokyo, 1989).

The invention also relates to antibodies themselves formed against the PepN protein or polypeptides derived thereof according to the invention, more particularly by immunization of appropriate animals with said polypeptides and recovery of antibodies thus formed.

It goes without saying that this production is not limited to polyclonal antibodies.

It also relates to any monoclonal antibody produced by any hybridoma liable to be formed according to classical methods from splenic cells of an animal, particularly of a mouse or a rat, immunized against the purified polypeptide of the invention on the one hand, and of cells of a myeloma cell line on the other hand, and to be selected by its ability to produce the monoclonal antibodies recognizing the polypeptide which has been initially used for the immunization of the animals.

The invention also relates to any antibody of the invention labelled by an appropriate label of the enzymatic, fluorescent or radioactive type.

The invention also relates to the use of such antibodies, for example for the detection of the PepN protein or polypeptides derived thereof, as described above.

The invention also relates to a process for preparing the nucleic acids according to the invention.

A suitable method for chemically preparing the single-stranded nucleic acids (containing at most 100 nucleotides of the invention) can be carried out according to the automatic β -cyanoethyl phosphoramidite method of DNA synthesis described in Bioorganic Chemistry 4; 274-325 (1986).

In the case of single-stranded DNA, the material which is obtained at the end of the DNA synthesis can be used as such.

A suitable method for chemically preparing the double-stranded nucleic acids (containing at most 100 bp of the invention) comprises the following steps:

- DNA synthesis of one sense oligonucleotide using the automatic b-cyanoethyl phosphoramidite method above cited, and DNA synthesis of one antisense oligonucleotide using either the above-mentioned automatic b-cyanoethyl phosphoramidite method, or enzymatic transcription of the sense-strand using a specific primer hybridizing to the 3'-end of the sense strand,
- combining the sense and antisense oligonucleotide by hybridization in order to form a DNA duplex,
- cloning the DNA duplex obtained into a suitable plasmid vector and recovery of the DNA according to classical methods such as restriction enzyme digestion and agarose electrophoresis, or by PCR amplification according to the procedure outlined above.

A method for the chemical preparation of nucleic acids with lengths greater than 100 nucleotides - or base pairs, in the case of double-stranded nucleic acids - comprises the following steps:

- assembling the synthesized oligonucleotides, provided at their ends with different restriction sites, the sequences of which are compatible with the succession of amino acids in the natural peptide, according to the principle described by Urdea et al. in Proc. Nat. Acad. Sci. USA 80; 7461-7465 (1983),
- cloning the DNA thereby obtained into a suitable plasmid vector and recovery of the desired nucleic acid according to classical methods such as restriction enzyme digestion and agarose gel electrophoresis.

Other characteristics and advantages of the invention will appear in the following examples and the figures illustrating the invention.

In cell extracts of *Lactobacillus delbrückii* ssp. *lactis* DSM7290 a peptidase with the activity to hydrolyse phe-β-naphthylamide and his-β-naphthylamide could be detected. *Escherichia coli* lacking the enzyme activity in a enzymatic plate assay, which was based on the hydrolysis of these β-naphthylamide (β-NA) substrates, was used to screen high copy and low copy number plasmid libraries of size fractionated *Lactobacillus* DNA. Clones with the desired phenotype were detected, and the gene, designated *pepN*, was further subcloned and

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sequenced. A large open reading frame of 2529 nt is predicted to encode a protein of 843 amino acids (95358 Da). Analysis of the *pepN* sequence indicated that the enzyme is not subjected to posttranslational modification or exported via processing of a signal peptide. Comparison of the *pepN* gene from *Lb. delbrückii* ssp. *lactis* DSM7290 indicates that it is homologous to genes of the family of Zn²+-metallohydrolases and shows identity with the active centre Zn²+-binding motif of these enzymes. Like in all other organisms the substrate lys-β-NA is more effectively cleaved than phe- or his-β-NA's, used for screening in *E. coli*. The cloned enzyme was extremely overexpressed in *E. coli* and subclonig of the gene in *Lactobacillus casei* resulted in a moderate overexpression of approximately 20-fold. The cloned enzyme was purified from the *pepN* deficient *E. coli* strain CM89, using the substrate lys-p-nitroanilide. In a four step procedure including streptomycin sulfate precipitation, anion exchange chromatography, and gelfiltration the peptidase was purified to electrophoretic homogeneity.

The structural proteinase genes (prtP: Kok et al., 1988; Kiwaki et al., 1989; Vos et al., 1989a; prtM: Haan-drickman et al., 1989; Kiwaki et al., 1989; Vos et al., 1989b;), the cysteine aminopeptidase (pepC, Chapot-Chartier et al., 1993), the X-prolyl-dipeptidyl-aminopeptidase genes (pepX; Nardi et al., 1991, Mayo et al., 1991), and the general aminopeptidase pepN (Stroman, 1992; van Alen-Boerrigter et al., 1991;) of a number of lactococci strains have been cloned and sequenced. From Lactobacillus strains only the proteinase genes (Holck and Naes, 1992). From Lb. paracasei and the X-prolyl-dipeptidyl-aminopeptidase gene from Lb. delbrūckii ssp. lactis DSM7290 (Meyer et al., 1993), have been cloned and sequenced. Lb. delbrūckii ssp. lactis is used as starter culture in the fermentation of swiss type cheese. Strain DSM7290, originally designated WS87, isolated from Emmenthaler cheese was chosen for screening of peptidase genes.

It was screened for complementation and expression in *E. coli*, with subtrates cleaved in cell extracts of DSM7290, but reacting negative in the enzyme plate assay used for screening. It has been decided to screen for heterologous expression in *E. coli* because of still moderate transformation efficiencies in lactobacilli and since peptidase activities in lactobacillal colonies cannot be reliably detected as a reason of acidification of agar plates and rigidity of the cell envelope.

I) Material and methods

Bacterial Strains, Plasmids, and Growth Conditions

The bacterial strains and plasmids used are summarized in Table 1. *Escherichia coli* was grown at 37°C in Luria broth (Miller, 1972), and lactobacilli in MRS (De Man et al., 1960) at 37°C. Ampicillin and kanamycin were added to concentrations of 100 and 40 µg/ml, respectively.

35 Transformations

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Lb. casei LK1 (Zink et al., 1991) and E. coli cells (Dower et al., 1988) were transformed by electroporation using a Bio-Rad Gene Pulser (Bio-Rad Laboratories, Richmond, CA) as described before.

40 Recombinant DNA Techniques.

Restriction enzymes and other nucleic acid-modifying enzymes were used as recommended by the manufacturers. Isolation of plasmid DNA from *E. coli* was performed as described by Sambrock et al. (1989). Plasmid DNA isolation from *Lb. casei* or *Lb. curvatus* was performed as described elsewhere (Zink et al., 1991).

Isolation of chromosomal DNA from DSM7290

400 ml of prewarmed MRS medium was inoculated with 40 ml of an overnight culture of *Lb. delbrūckii* ssp. *lactis* DSM7290 and incubated for 2.5 hours at 37°C. Cells were pelleted by centrifugation, washed once with 50 mM Tris-HCl pH 8, resuspended in 37,5 ml of sucrose 6.5 %, 50 mM Tris-HCl pH 8, 1 mM EDTA, 200 mg of Lysozym, and 2500 U of mutanolysin (Sigma) and incubated for 1 hour at 37°C. 3.75 ml of 0.25 M EDTA in 50 mM Tris-HCl pH 8 was added. Cells were lysed by the addition of 2.25 ml 20 % SDS. The lysed cells were subjected to proteinase K digestion at a concentration of 50 μ g/ml, for 15 min at 50°C and 30 min at 60°C. EtBr was added to a concentration of 1 mg/ml and CsCl to reach a final densitity of 1.55 g/ml. Centrifugation to equilibrium and further purification of chromosomal DNA was performed as described by Sambrock et al. (1989).

Molecular cloning of the Lb. delbrūckii ssp. lactis pepN gene

Construction of plasmid libraries:

Genomic plasmid libraries of *Lb. delbrūckii* ssp. *lactis* DSM7290 were prepared in *E. coli* ER1562 using the positive selection vector pUH84 (Henrich and Plapp, 1984) and because overexpression of peptidase genes from this high copy number vector may be lethal to the host a second plasmid bank was constructed with vector pLG339 having only 6-8 copies per chromosome (Stoker et al., 1982). Partial *Sau*3A fragments of total DSM7290 DNA were size fractionated (10-40 % sucrose gradient) and 4-10 kb fragments were ligated into the dephosphorylated *Bam*HI sites of the vector molecules.

Identification of peptidase genes:

Colonies of *E. coli* transformed with the gene banks were screened for PepN activity by a plate staining method of Miller and Mackinnon (1974), recently used by Nardi et al. (1991). If chromogenic β -NA substrates are cleaved, reaction of the β -napthylamines with fast garnet CBC (Sigma), can be monitored by the formation of a red, non-diffusible azo dye.

DNA Sequence Analysis

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For nucleotide sequencing of inserts in pLG339, a pair of universal sequencing primers, adjacent to the *Bam*HI cloning site was synthesized. Synthetic oligonucleotide primers deduced from the investigated sequence, were synthesized and allowed direct sequencing of double stranded plasmid DNA. The DNA sequence of each strand was determined using the T7 DNA polymerase sequencing kit (Pharmacia, Uppsala, Sweden), which is based on the dideoxynucleotide chain termination method (Sanger et al., 1977) in the presence of $[\alpha^{-35}S]$ dATP (Amersham). For computer-assisted sequence analysis the Microgenie (Beckman, Palo Alto, CA), PC-Gene (IntelliGenetics, East El Camino Real, CA), and HUSAR (GENIUSnet, Heidelberg) software were used.

Sequencing primers were synthesized using Applied Biosystems Model 392 DNA synthesizer and reagents.

Preparation of cell extracts

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Cell pellets of *E. coli* from a 2 I overnight culture were washed with 50 mM Tris-HCI pH 8.0, pelleted by centrifugation and resuspended in 20 ml of the same buffer. The bacteria were sonicated on ice (Bandelin sonifier; Sonopuls HD60) until more than 90 % of the cells were broken. Cell debris were removed by centrifugation at 52,000 g and 4°C for 60 min.

Lactobacillal cell extracts were prepared with 50 ml cultures and the modification that prior sonification of the more rigid bacteria, glass beads (diameter, 0.17-0.18 mm) were added to the cell pellets (2 Vol glass-beads/ 1 Vol cells). Centrifugation was performed in a Hereus Biofuge RS28 at 51,000 g and 4°C for 60 min.

The supernatants of both preparations contained approximately 50 mg/ml of protein as determined by the method of Lowry (1951).

Enzyme assay and effects of various chemical reagents in crude cell extracts

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For characterization of the enzyme the p-nitroanilide subtrates (Bachem) were dissolved in water and added to the reaction mixture [Tris-HCl 10 mM, pH 8.0, and varying amounts of purified enzyme (1-200 ng)] to a concentration of 1 mM in a volume of 250 µl. Release of p-nitroaniline was measured after a 10 min incubation at 37°C at 405 mm in a LKB Ultrospec plus spectrophotometer. For rapid screening of active fractions during purification of the enzyme, samples were prepared in micro titer plates and measured in a BIORAD model 2550 EIA reader.

To study the mechanism of enzyme action, the inhibitors phenylmethan sulphonyl fluoride (PMSF) at a concentration of 1 mM, pepstatin A at 1 µg/ml, L-trans-epoxy succinyl-leucylamide(4-guanidino)-butane (E-64) at 0.1 mM, and 1,10-phen anthroline or EDTA at 1 mM were added to the enzyme and incubated for 30 min at 37°C. The substrate lys-p-nitro anilide was added and activity was measured spectrophotometrically by the release of p-nitro aniline.

Isoelectric focusing

Isoelectric focusing was carried out in a LKB column (type 8100-1) with ampholytes (Serva) ranging from pH 4 to 6. The pH gradient was stabilized by a glycerol gradient.

Purification

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Step 1: Streptomycin sulfate precipitation

At 4°C a solution of 10 % streptomycin sulfate was slowly added to cell extract while stirring, to reach a final concentration of 2 %. The precipitate was discarded after centrifugation (10 min, 20,000 g.)

Steps 2 and 3: Anion exchange chromatography

The supernatant of the streptomycin sulfate precipitation was concentrated by passage through a PM30 Diaflow membrane (Amicon) and applied to a 38 by 2.2 cm column of Q-sepharose fast flow (Pharmacia) equilibrated with 0.1 M NaCl in 20 mM Tris-HCl pH 8.0 at 4°C. Proteins were eluted with a linear NaCl gradient of 0 to 0.5 M NaCl respectively in the same buffer. In a second passage a gradient ranging form 0.4 to 0.7 M NaCl was chosen. 4 ml fractions were collected at a flow rate of 150 ml/h and those with high PepN activity were pooled and concentrated by passage through a PM30 Diaflow membrane.

Step 4: Gelfiltration.

The enriched PepN preparation from anion exchange chromatography was applied to a 55 by 2.2 cm column of Fractogel TSK HW-55(S) (Merck) equilibrated with 20 mM Tris-HCl pH 7.2 at 4°C. Elution was performed with the same buffer at 20 ml/min and 2 ml fractions were collected.

II) Results and Discussion

30 Cloning of the Lb. delbrückii ssp. lactis DSM7290 pepN gene.

The plasmid libraries of DSM7290 chromosomal DNA in pUH84 and pLG339 respectively were transformed into *E. coli* ER1562. The *pepN* gene could be isolated by screening the colonies for peptolytic activities using the chromogenic substrates his-β-NA and phe-β-NA. In the plate assay these substrates were not cleaved by the homologous PepN of the *E. coli* host strain, and additionally they were not cleaved by *E. coli* harbouring the recently cloned genes for *pepX* (X-prolyl-dipeptidyl aminopeptidase) (Meyer et al., 1993) and *pepP* (prolin iminopeptidase, unpublished) from *Lb. delbrūckii* ssp. *lactis* DSM7290. The gene has been designated *pepN* since sequence analysis revealed extensive amino acid homology to other aminopeptidases N (E.C. 3.4.11.2).

From approximately 50,000 colonies harbouring high copy number genebank plasmids no positive reaction could be detected in the plate assay, but in transformants with the low copy number plasmid bank, with pLG339 as vector, 3 his- β -NA and 3 phe- β -NA cleaving colonies out of 21,000 could be detected. The inability to identify positive clones in the pUH84 genebank indicated that overexpression of the peptidase gene is lethal for *E. coli*. Restriction analysis revealed that all plasmids isolated had insert sizes ranging from 5.2 to 8.0 kb, with a conserved identical core region. This agreed with the observation that all plasmids coded for an enzyme cleaving both substrates his- β -NA and phe- β -NA.

The plasmid pJUK11 with the smallest insert size of 5.2 kb was subject for further experiments (FIG.1). The chromogenic β-NA substrates have been used for colony screening and the analogous p-nitroanilides for characterization of the enzyme.

Plasmid encoded proteins

E. coli cells harbouring pJUK11 were sonified and the crude cell extracts subjected to SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis) (Laemmli, 1970). A major 95 kDa protein appeared in ER-1562(pJUK11) extracts which constituted at least 50 % of the cytoplasmic E. coli proteins (FIG.2). This remarkable high level expression of pepN might be the reason for its lethality if cloned in a high copy number plasmid.

Subcloning and nucleotide sequence analysis

Prior to sequence analysis the region coding for *pepN* was localized, calculating that approximately 2.6 kb should be sufficient to code for the 95 kDa protein detected by SDS-PAGE. Subcloning of a 3.2 kb *Sphl/Hind*III fragment from pJUK11 in pUC18 (FIG.1) resulted in a plasmid coding for the *pepN* gene with expression of enzyme activity, but which was extremely unstable. Nevertheless it offered the possibility to determine the DNA sequence starting at the HindIII site with the universal primer of vector pUC18. The second strand, which is coding for *pepN* was sequenced using a synthesized 20mer primer, complementary to the tetracyclin resistance gene in pJUK11. The complete nucleotide sequence of both strands of the *Sphl/Hind*III fragment was determined using synthetic oligonucleotide primers deduced from the investigated sequences. The lactobacillal nucleotide sequence determined (3122 bp, 49.4% C+G) contained one single open reading frame (FIG.3). Starting with a putative ATG codon it would code for 843 amino acids corresponding to a protein with a molecular weight of 95358 Da. This value corresponds well with the molecular weight of the enzyme predicted by SDS-PAGE.

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Transcription of the pepN gene

Upstream of the ATG start codon spaced by six base pairs, a putative ribosome binding site with a hexanucleotide stretch (AGGAGG) complementary to the *E. coli* 16 S-rRNA, and sequences that resemble promoter -10 and -35 sequences (Fig. 3) elements of gram positives (Van de Guchte et al., 1992) can be located. Currently we are going to compare the functionality of these sequences in *E. coli* and *Lb. delbrückii* ssp. *lactis*. The *pepN* coding sequence is followed by an inverted repeat which may form a stem-loop structure with a dG of -12.8 kcal/mol, and which may function as a transcriptional terminator, as predicted by the computer algorithm of Brendel and Trifonov (1984).

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Comparison of the lactobacillal PepN sequence to aminopeptidases of other species

Searching the EMBL database resulted in similarities to several lys-aminopeptidases (FIG.4) belonging to the aminopeptidase N family (E.C. 3.4.11.2). These enzymes, sharing a common pattern of primary structure in the part of their sequence involved in the binding of zinc, can be grouped together as a family of neutral zinc metallopeptidases. They catalyse the removal of N-terminal amino acids from peptides. The most significant homology is observed in a functionally important segment which contains the proposed Zn²+-binding catalytic site (Jongeneel et al., 1989). The phylogenetic tree calculated (FIG.5) demonstrates the close relationship of the lactobacillal enzyme to those of the lactococcal species. The diversity in the N-terminal sequences is a result from signal sequences present in human, rat, and mouse enzymes. The absence of a signal peptide sequence and the hydrophilicity plot of the amino acid sequence according to Kyte and Doolittle (1982) which does not show transmembrane domains, indicate that PepN of *Lb. delbrūckii* ssp. *lactis* might be a intercellular located enzyme. Until now, no structure allowing protein secretion is described for any peptidase of lactic acid bacteria, but since the action of peptidases is required for casein cleavage a yet unknown mechanism might be responsible for translocation to the cell surface.

Gene dosis effects

The knowledge of the *pepN* nucleotide sequence allowed subcloning of a minimal sized DNA fragment into different cloning vectors. As summarized in FIG.1 we succeeded in cloning a 2.9 kb Dral fragment of pJUK11 in both orientations into pLG339, thus indicating that the *pepN* promoter sequence is functional in *E. coli*. This fragment was also cloned into the medium copy number plasmid pBR322 and even into the high copy number plasmid pUC18. In the latter constructs only one of the two possible insert orientations were obtained, likely an effect of lethal overproduction, since *pepN* would then be under control of lac or tet promoters. The amount of PepN protein produced, increased with the copy number of the plasmids, but on costs of plasmid stability (FIG.6).

For homologous expression in a *Lactobacillus* starter culture the cloned peptidase gene was inserted in both orientations, as a *Sph*I-restriction fragment, into pJK355 (FIG.1), a *Lactobacillus* vector constructed from the cryptic plasmid from *Lb. curvatus* LTH683 (Klein et al., 1993) resulting in plasmids pJK361 and pJK362. In crude extracts of *Lb. casei* LK1, with a chromosomal *pepN* gene, a lys-pNA hydrolysing activity could be measured, but expression of *pepN* coded on plasmids pJK361 or pJK362 resulted in overproduction of enzyme visible in SDS-PAGE (FIG.3) and enzymatic activities in crude extracts were increased by 15 to 20 fold. This increase of PepN activity had no obvious effect on cell growth in MRS-broth, which could have been expected

due to an imbalance in peptide supply. The influence on growth in milk is still going to be investigated as well as the effects on changes in taste and texture of cheese produced with such a modified strain. Purified PepN of *Lc. lactis* is reported to have a debittering activity on tryptic digests of β -casein (Tan et al., 1992). If the cloned and overexpressed enzyme of DSM7290 has comparable activities this gene can be of great industrial interest.

Purification and partial characterization of the cloned peptidase N

Since PepN protein was extremely overexpressed in *E. coli* harbouring the plasmid coded gene, the enzyme could be purified to electrophoretic purity by use of a four-step procedure. Due to sequence homologies of the cloned lactobacillal *pepN* with the *Escherichia coli* aminopeptidase N gene, the enzymatic properties were investigated in CM89 a strain lacking the *E. coli* enzyme. CM89(pUK13) was chosen for purification because the plasmid was stable in the presence of Kanamycin and high yields of PepN protein were obtained. Table 2 summarizes the purification to electrophoretic purity by the use of streptomycin sulfate precipitation, anion exchange chromatographies on Q-sepharose, and gelfiltration on HW TSK 55(S). The final purification was approximately 11-fold with a recovery of 20 %. Estimated by SDS-PAGE and gelfiltration, the enzyme is a monomer with a molecular mass of 95 kDa, which agrees well with the value deduced from sequence analysis. The pl of 4.2, determined by isoelectric focusing agrees well with the predicted one of 4.48 deduced from amino acid sequence. Optimum PepN activity was measured at pH of 6.5 - 7.0 and temperatures of 45 - 55°C. Denaturation occurs at pH values below 4 and at temperatures above 60°C.

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The classification based on the susceptibility to a group of protease inhibitors (PMSF, E-64, pepstatin, EDTA, and 1,10 phenanthroline) indicated that the enzyme belongs to the class of metallo proteases, as full inhibition with 1,10 phenanthroline or EDTA could be detected, whereas the inhibitors PMSF, Pepstatin A, or E-64 had no effect on enzyme activity. Additionally the homology to other aminopeptidases implies that PepN has a catalytic centre with zinc binding sites (FIG.4). The inhibition by both complexing agents could be specifically restored by addition of Mn²⁺ and Co²⁺, while Zn²⁺ or Mg²⁺ had no effect. Therefore one might suspect that Mn²⁺ and Co²⁺ ions play an essential role in the hydrolytic mechanism.

The substrate specificity of PepN against various p-nitroanilides is shown in Table 3. The highest activity was found for lys-p-nitroanilide and a Lineweaver-Burk plot indicated a Km of 70 µM. In comparison with the enzyme of *Lc. lactis* ssp. *cremoris* (Tan and Konings, 1990), showing the closest sequence relationship (62 % similarity) the lactobacillal Km value demonstrates an 8-fold higher affinity against lys-pNA. There are certain similar properties such as specifity against several p-NA substrates with bulk or aliphatic N-terminal residues, and pH optimum. The enzyme differs in restoration of enzyme activity after 1,10 phenanthroline inhibition. The temperature optimum of 50°C and its stability at elevated temperatures, after a 20 min incubation at 60°C a residual activity of 33 % were measured, might be of interest for applications of the cloned enzyme in milk fermentations.

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Legends to the figures

FIG. 1. Circular map of plasmid pJUK11. The insert of chromosomal DNA of 5.2 kbp in size, is indicated as a shaded segment, with the *pepN* gene presented as dotted arrow. The linear expansion of the insert illustrates the sequencing strategy. The different restriction fragments used for subcloning into different vectors, all spanning the *pepN* gene, are indicated as black bars.

FIG. 2. Expression of PepN protein in *E. coli* (ER1562) and *Lb. casei* (LK1). Separation of crude cell extracts by SDS-PAGE (12 % acrylamide). Proteins were visualized by Coomassie staining. Lanes 1 and 2, molecular weight marker proteins; cell extracts of *E. coli* ER1562(pJUK11) [lane3] and ER1562(pLG339) [lane4]; cell extracts of LK1 [lane 7], LK1(pJK361) [lane5] and LK1(pJK362) [lane 6]. The position of PepN is marked by an arrow head.

FIG. 3. Nucleotide sequence of the *pepN* region and deduced amino acid sequence of PepN. The proposed -35 (212 to 234) and -10 (250 to 255) positions and ribosome binding site (305 to 309) are underlined. An inverted repeat (underlined at bps 2871 to 2891) with a dG value of -12.8 kcal/mol, followed by a run of T-residues, is proposed as a transcription terminator.

FIG. 4. Progressive alignment (TREE, software of HUSAR GENIUSnet, Heidelberg) of the protein sequence of PepN from *Lb. delbrūckii* ssp. *lactis* DSM7290 with those from other aminopeptidases. Identical amino acids in the sequences are marked (*). The putative catalytic sites (A) with zinc-binding domains (Z) are indicated.

FIG. 5. Phylogenetic relationship of aminopeptidases from different species. The dendrogram was calcu-

lated from the progressive alignment of homologous protein sequences using the program TREE of HUSAR, which is based on the method described by Feng and Doolittle (1987). The vertical bars represent nucleotide changes in percent.

FIG. 6. Stability of different plasmids coding for *pepN* in *Escherichia coli*. ER1562 harbouring different plasmids (pJUK12, pJUK13, pJUK14, pJUK15) was grown in liquid medium with and without the appropriate antibiotic selection.

Table 1. Bacterial strains and plasmids

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Table 3. Relative activity of PepN for p-nitroanilide substrates

p-nitroanilide (-pNA) substrate	relative activity (%)
Lys-pna	100
Leu-pNA	19
Ala-pNA	12
Phe-pNA	8 .
Pro-pNA	2
Tyr-pNA	2
Gly-pNA	1
Ala-Pro-pNA	0

SEQUENCE LISTING

5	
	(1) GENERAL INFORMATION:
	(i) APPLICANT:
	(A) NAME: COMMUNAUTE ECONOMIQUE EUROPEENNE
10	(B) STREET: BATIMENT JEAN MONNET - PLATEAU DU KIRCHBERG
	(C) CITY: LUXEMBOURG
	(D) STATE: LUXEMBOURG
	(E) COUNTRY: LUXEMBOURG
	(F) POSTAL CODE (ZIP): L2920
15	(ii) TITLE OF INVENTION: LYS-AMINOPEPTIDASE PepN FROM LACTOBACILLUS
	DELBRUCKII SSP. LACTIS, NUCLEIC ACIDS CODING FOR IT, AND
	ITS USE IN FERMENTATION PROCESSES
	• •
	(iii) NUMBER OF SEQUENCES: 2
20	(iv) COMPANIES DESCRIPTION
	(iv) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk
	(B) COMPUTER: IBM PC compatible
	(C) OPERATING SYSTEM: PC-DOS/MS-DOS
25	(D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
25	·
	(vi) PRIOR APPLICATION DATA:
	(A) APPLICATION NUMBER: GB 93-13586.1
	(B) FILING DATE: 01-JUL-1993
30	
	(2) INFORMATION FOR SEQ ID NO: 1:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 3122 base pairs
35	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: double (D) TOPOLOGY: linear
	(b) lorobodi: lillear
	(ii) MOLECULE TYPE: DNA (genomic)
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40	(ix) FEATURE:
	(A) NAME/KEY: CDS
	(B) LOCATION: 3162844
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
45	(, Degree Beschifficht. Beg ID NO: 1:
	GATAGGGCTG AAATTATCAT TTTAAGCGCT TTAAATTAGC TATACAGATA AGTAACATTA 60
	GTAACAATTG TCAAGAGACT GCAATAAAAG GAAAAGGCCA GCTGCTAGAC TGGTCTTTTA 120
	CATATGCAAT TATTTCAAAA AMGGAAMMAA MUUGAAMMAA
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	GTTCAAGGGG CTGAGGTAGA CTGCAAACAG ATATTTTGCG TTAAATGGGC TTTATTTAGC 240
	CTTTTTTGCT AGAATAGAGA AGTGTGAATA CAATATACGC GAGGAAAAAT CAGACGCGGA 300
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			Thr									Glu			TTG Leu	447
15		His	CAA Gln			Thr								•	AAG Lys	495
20			CCG Pro		Phe										GAA Glu	543
25			AAG Lys													591
20			GAC Asp 95													639
30			AAG Lys													687
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			CCA Pro													831
45			CGC Arg 175													879
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						TGG Trp													1119	
	15					AAC Asn													1167	
		GTG	GTG	ACC	CAC	GAA	CTG	GCC	CAC	CAA	TGG	TTC	GGT	GAC	CTG	СТЪ	ACC		1215	
	20					Glu														
v.						GAC Asp 305												;	1263	
	25																			
		ATG Met	GAA Glu	TAC Tyr	CTG Leu 320	TCA Ser	GTT Val	GAC Asp	CAC His	CTG Leu 325	GAA Glu	CCT Pro	AAC Asn	TGG Trp	CAC His 330	ATC Ile	TGG Trp	;	1311	
	30					ACT Thr												:	1359	
	35					CAG Gln												:	1407	
		ATC Ile 365	GAC Asp	GCC Ala	CTC Leu	TTT Phe	GAC Asp 370	GGG Gly	GCC Ala	ATC Ile	GTT Val	TAC Tyr 375	GCC Ala	AAG Lys	GGG Gly	TCA Ser	AGA Arg 380	:	1455	
	40	ATG Met	CTG Leu	GTC Val	ATG Met	GTC Val 385	CGG Arg	TCA Ser	CTT Leu	Leu	GGC Gly 390	GAT Asp	GAA Glu	GCC Ala	TTG Leu	AGA Arg 395	AAG Lys	=	1503	
	45	GGC Gly	TTG Leu	AAG Lys	CGC Arg 400	TAC Tyr	TTT Phe	GAC Asp	Lys	His 405	AAG Lys	TTT Phe	GGC Gly	AAC Asn	GCG Ala 410	GCA Ala	GGT Gly	. 1	1551	
, 	50	GAC Asp	Asp	CTC Leu 415	TGG Trp	GAT Asp	GCC Ala	Leu	TCA Ser 420	ACG	GCC Ala	ACT Thr	GAC Asp	TTG Leu 425	AAC Asn	ATT Ile	GGG Gly	1	L599	16
		GAA Glu	ATC Ile 430	ATG Met	CAC . His	ACT '	Trp	CTG Leu 435	GAC Asp	CAG Gln	CCA Pro	Gly	TAC Tyr 440	CCA Pro	GTG Val	GTG Val	AAT Asn	1	1647	

5	_	Phe			GAC Asp	His			Gln				1695
	_				AAG Lys 465			Trp			-	AAC Asn	1743
10					GCA Ala								1791
15				Gln	GCT Ala								1839
20			Gly		AAC Asn								1887
					ATG Met								1935
25					CAA Gln 545								1983
30					GTG Val								2031
35					AAC Asn								2079
					GCC Ala								2127
40					TCC Ser								2175
45					AGC Ser 625								2223
50					TAC Tyr								2271
					GAA Glu								2319

		ATC	CGG	CCA	TAC	GTT	TTG	ATC	AAC	GAA	GTT	GAA	AAC	TAC	GGG	TCA	AGC	2367		
			Arg		Tyr			Ile					Asn							
	5		670					675					680							
		_			GAC													2415		
		685	Leu	Thr	Asp	Lys	Leu 690	Ile	Gly	Leu	Tyr	Gln 695	Ala	Thr	Ser	Asp	Pro 700			
																	700			_
	10				ATG													2463		
		ser	Pne	гуз	Met	705	Leu	GIU	Ата	АТА	710	vaı	Lys	Ser	Lys	715	Glu			
							•													
	15				AAG													2511		
:	10	GIY	GIU	пец	Lys 720	пуз	me	val	ser	725	Pne	гув	ASI	Ата	730	TIE	vai			
																•				
				_	GAC Asp													2559		
	20	2,0	110	735	лор	пси	ar 9	Cly	740	rne	Der	Gry	Val	745	SEL	ASII	PIO	•		
		~~~		~																
					CAG Gln													2607		
•			750					755					760	014	11.0		115			
•	25	mma	<b>(13.3</b>	220	200	ama.	999	<b>666</b>	~~~	3 ma	<i>a</i>									
					ACG Thr													2655		
		765		-			770	•				775					780			
		GTC	ATC'	<b>ייי</b> רירי	CGC	GTC.	ጥጥር	מממ	NCC.	220	CAA	~~~	TTA C	CAC	<i>(</i> (2) 2)	ma.c	220	2702		
	30				Arg													2703		
						785					790		_	_		795				
		GCC	TTC	TTT	ACT	GAC	AAG	GAA	AGC	AAC	ATG	CTG	СТС	AAC	rcc	GDD	<b>ል</b> ጥር	2751		
					Thr													2751		
	35				800					805					810				•	
		AAG	ATG	GAC	CGG	AAG	GTC	ATC	GCT	AAC	CGG	GTA	GAC	TTG	ATT	GCC	AGC	2799		
				Asp	Arg															
				815					820					825						
	40	GAA	CAA	GCT	GAC	GTC	AAC	GCC	GCG	GTT	GCT	GCT	GCT	TTG	CAA	AAG		2844		
		Glu		Ala	Asp	Val	Asn		Ala	٧al	Ala	Ala		Leu	Gln	Lys				
			830					835					840							
	45	TAAT	'TGAA	TA G	BAGCA	TAAG	A AA	ACTO	TTTC	CGC	TGAG	AGC	TGG	AACA	GT I	TTTT	TATGT	2904		
	43	ልጥጥር	ים ממי	ייזיכי יו	יכיייכיר	יה אידיר	'C CT	יידיא <i>ר</i> יזי	אייי איי	י אריא	מייטים.	אסת	a curr	maan	120 2	mmon	TTGAG	2054		
(()	-	AIIC	AMCI	.16 1	·	AAIC	.6 61	IACA		AGA	TGTA	LAAT	ACTA	rrcgi	AC A	TTC1	TTGAG	2964		
		GTAA	TAAA	AT G	AACA	ACGA	T TI	TAAA	GATA	TCA	TGCA	AAA	CAGA	AAGT	CT A	TCC	GCACT	3024		
	50	ATGA	TTCC	AG C	GTGA	AGAT	יד דר	CCGT	'GACG	דמת :	ייזינגרייז	'GC'A	ስምር	מידיים <b>י</b>	י ייע ב	! ል ውጥር	TATCT	3084		
													WHI.	·WT TW		mara I (	IMICI	3004		
		CTGC	TCCA	AG I	GCCI	GCAA	C CT	GCAG	TCCI	GGA	AGCI	T						3122		

(2) INFORMATION FOR SEQ ID NO: 2:

5			()	SEQU A) L B) T D) T	ENGT YPE :	H: 8	43 a	mino cid								
		(ii	) MO	LECU	LE T	YPE:	pro	tein								
10		(xi	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0: 2	:				
	Met 1	Ala	Val	Lys	Arg 5	Phe	Tyr	Glu	Thr	Phe 10	His	Pro	Asp	His	Tyr 15	Asp
15	Leu	Tyr	Ile	Asp 20	Val	Asp	Arg	Ala	Ala 25	Arg	Ser	Phe	Ser	Gly 30	Thr	Ser
20	Thr	Ile	His 35	Gly	Glu	Ile	Gln	Glu 40	Glu	Thr	Val	Leu	Val 45	His	Gln	Lys
	Tyr	Met 50	Thr	Ile	Ser	Lys	Val 55	Thr	Val	Asp	Gly	60	Glu	Val	Pro	Phe
25	Thr 65	Phe	Gly	Asp	Asp	Phe 70	Glu	Gly	Ile	Lys	Ile 75	Glu	Ala	Gly	Lys	Thr 80
			Ala	Val	Ile 85	Ala	Ile	Asp	Tyr	Ser 90	Ala	Pro	Leu	Thr	Asp 95	Thr
30		Met	Gly	Ile 100	Tyr	Pro	Ser	Tyr	Tyr 105	Gln	Val	Asp	Gly	Val 110	ГÀЗ	ГÀЗ
35	Glu	Leu	Ile 115	Gly	Thr	Gln	Phe	Glu 120	Thr	Thr	Phe	Ala	Arg 125	Glu	Ala	Phe
	Pro	Cys 130	Val	Asp	Glu	Pro	Glu 135	Ala	Lys	Ala	Thr	Phe 140	Ser	Leu	Ala	Leu
40	Lys 145	Phe	Asp	Glu	His	Glu 150	Gly	Glu	Thr	Val	Leu 155	Ala	Asn	Met	Pro	Glu 160
	Asp	Arg	Val	Glu	Asn 165	Gly	Val	His	Tyr	Phe 170	Lys	Glu	Thr	Val	Arg 175	Met
45	Ser	Ser	Tyr	Leu 180	Val	Ala	Phe	Ala	Phe 185	Gly	Glu	Met	Arg	Ser 190	Leu	Thr
<b>5</b> 0	Thr	His	Thr 195	Lys	Ser	Gly	Val	Leu 200	Ile	Gly	Val	Tyr	Ser 205	Thr	Gln	Ala
50	His	Thr 210	Glu	Lys	Glu	Leu	Thr 215	Phe	Ser	Leu	Asp	Ile 220	Ala	Lys	Arg	Ala
55	Ile 225	Glu	Phe	Tyr	Glu	Asp 230	Phe	Tyr	Gln	Thr	Pro 235	Tyr	Pro	Leu	Pro	Gln 240
	Ser	Leu	Gln	Leu	Ala 245	Leu	Pro	Asp	Phe	Ser	Ala	Gly	Ala	Met	Glu 255	Asn

		1	p Gi	y ne	260	. Thi	r Ty:	r Arg	g GT	u A1 26		r Le	u Le	u Lei	u As; 27		o Asp
	5	Ası	n Thi	r Thi 275	r Leu 5	Glu	ı Mei	t Lys	280	s Le O	u Vai	l Ala	a Th:	r Val 28!		l Th	r His
		Glı	290	ı Ala	A His	Glr	Tr	Phe 295	e Gly	y As	p Lei	ı Va	1 Th:		Gl:	u Trj	p Trp
	10	Asp 305	o Ası 5	ı Lev	ı Trp	Leu	Asr 310	ı Glı )	ı Sei	r Ph	e Ala	31:		: Met	Glı	1 Ту	r Leu 320
	15	Ser	· Val	. Asp	His	Leu 325	Glu	ı Pro	) Ası	ı Tr	9 His		Tr	Glu	ı Met	2 Phe 335	e Gln 5
		Thr	Ser	Glu	Ala 340	Ala	Ala	Ala	Let	1 Th:		As <u>r</u>	Ala	a Thr	Asp 350		/ Val
	20	Gln	ser	Val 355	His	Val	Glu	Val	Asr 360	ı Ası	Pro	Ala	Glu	1 Ile 365		Ala	Leu
٠		Phe	370	Gly	Ala	Ile :	Val	Tyr 375		Lys	s Gly	Ser	Arg		Leu	ı Val	. Met
	25	Val 385	Arg	Ser	Leu	Leu	Gly 390	Asp	Glu	Ala	Leu	Arg 395		Gly	Leu	Lys	Arg 400
	<b>30</b> .	Tyr	Phe	Asp	Lys	His 405	Lys	Phe	Gly	' Asn	Ala 410		Gly	Asp	Asp	Leu 415	Trp
	••	Asp	Ala	Leu	Ser 420	Thr	Ala	Thr	Asp	Leu 425		Ile	Gly	Glu	Ile 430		His
	35	Thr	Trp	Leu 435	Asp	Gln	Pro	Gly	Tyr 440	Pro	Val	Val	Asn	Ala 445	Phe	Val	Glu
		Asp	Gly 450	His	Leu	Lys	Leu	Thr 455	Gln	Lys	Gln	Phe	Phe	Ile	Gly	Glu	Gly
	40	Lys 465	Glu	Val	Gly	Arg	Lys 470	Trp	Glu	Ile	Pro	Leu 475	Asn	Ala	Asn	Phe	Lys 480
		Ala	Pro	Lys	Ile	Met 485	Ser	Asp	Val	Glu	Leu 490	Asp	Leu	Gly	Asp	Tyr 495	Gln
	<b>.</b> -	Ala	Leu	Arg	Ala 500	Glu	Ala	Gly	His	Ala 505	Leu	Arg	Leu	Asn	Val 510	Gly	Asn
	50	Asn	Ser	His 515	Phe	Ile	Val	Lys	Tyr 520	Asp	Gln	Thr	Leu	Met 525	Asp	Asp	Ile
		Met	Lys 530	Glu	Ala :	Lys	Asp	Leu 535	Asp	Pro	Val	Ser	Gln 540	Leu	Gln	Leu	Leu
	55	Gln 545	Asp	Leu	Arg 1	Leu	Leu 550	Ala	Glu	Gly	Lys	Gln 555	Ala	Ser	Tyr	Ala	Asp

	Val	Val	Pro	Val	Leu 565	Glu	Leu	Phe	Lys	Asn 570	Ser	Glu	Ser	His	Ile 575	Val
5	Asn	Asp	Ala	Leu 580	Tyr	Thr	Thr	Ala	Asp 585	Lys	Leu	Arg	Gln	Phe 590	Ala	Pro
	Ala	Gly	Ser 595	Glu	Ala	Asp	Lys	Asn 600	Leu	Arg	Ala	Leu	Tyr 605	Asn	Asp	Leu
10	Ser	Lys 610	Asp	Gln	Val	Ala	Arg 615	Leu	Gly	Trp	Leu	Pro 620	Lys	Ala	Gly	Glu
15	Ser 625	Asp	Glu	Asp	Ile	Gln 630	Thr	Arg	Pro	Tyr	Val 635	Leu	Ser	Ala	Ser	Leu 640
	Tyr	Gly	Arg	Asn	Ala 645	Asp	Ser	Glu	Lys	Gln 650	Ala	His	Glu	Ile	Tyr 655	Val
20	Glu	Tyr	Ala	Asp 660	Lys	Leu	Ala	Glu	Leu 665	Ser	Ala	Asp	Ile	Arg 670	Pro	Tyr
	Val	Leu	Ile 675	Asn	Glu	Val	Glu	Asn 680	Tyr	Gly	Ser	Ser	Glu 685	Leu	Thr	Asp
25	Lys	Leu 690	Ile	Gly	Leu	Tyr	Gln 695	Ala	Thr	Ser	Asp	Pro 700	Ser	Phe	Lys	Met
30	Asp 705	Leu	Glu	Ala	Ala	Ile 710	Val	Lys	Ser	Lys	Asp 715	Glu	Gly	Glu	Leu	Lys 720
	Lys	Ile	Val	Ser	Trp 725	Phe	Lys	Asn	Ala	Glu 730	Ile	Val	Lys	Pro	Gln 735	Asp
35	Leu	Arg	Gly	Trp 740	Phe	Ser	Gly	Val	Leu 745	Ser	Asn	Pro	Ala	Gly 750	Glu	Gln
	Leu	Ala	Trp 755	Asp	Trp	Ile	Arg	Asp 760	Glu	Trp	Ala	Trp	Leu 765	Glu	Lys	Thr
40	Val	Gly 770	Gly	Asp	Met	Glu	Phe 775	Ala	Thr	Phe	Ile	Thr 780	Val	Ile	Ser	Arg
45	Val 785	Phe	Lys	Thr	Lys	Glu 790	Arg	Tyr	Asp	Glu	Tyr 795	Asn	Ala	Phe	Phe	Thr 800
45	Asp	Lys	Glu	Ser	Asn 805	Met	Leu	Leu	Asn	Arg 810	Glu	Ile	Lys	Met	Asp 815	Arg
50	Lys	Val	Ile	Ala 820	Asn	Arg	Val	Asp	Leu 825	Ile	Ala	Ser	Glu	Gln 830	Ala	Asp
	Val	Asn	Ala 835	Ala	Val	Ala	Ala	Ala 840	Leu	Gln	Lys				•	

#### Claims

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- Protein in a substantially pure form, as produced and excreted by Lactobacillus delbrūckii ssp. lactis and capable of hydrolysing aminoacid β-naphtylamides, and more particularly lysine-β-naphtylamide, or fragments thereof having this enzymatic activity.
  - Protein according to claim 1, such as produced by Lactobacillus delbrūckii ssp. lactis WS87, deposited at the Deutsche Sammlung von Mikroorganismen (DSM) under the number 7290 on October 15, 1992.
- 3. Protein according to claim 1 or claim 2, characterized in that it corresponds to the one (also called PepN) represented on figure 3, and by SEQ ID NO 2, or fragments or muteins (which differ from said protein by addition and/or substitution and/or suppression of one or several amino acid) thereof, provided that said fragments and muteins are capable of hydrolysing β-naphtylamides.
  - 4. Protein according to anyone of claims 1 to 3, characterized in that:
    - its isoelectric point calculated from the nucleotide sequence represented by SEQ ID NO 1, is 4,48, and its isoelectric point determined by preparative isoelectric focusing after purification of said protein, is 4,2,
    - its molecular weight calculated from the nucleotide sequence represented by SEQID NO 1, is 95,358
       kDa, and its molecular weight determined after purification, is 95 kDa,
    - its specific inhibitors are the following:
      - . EDTA: end concentration 1mM; 3 % relative activity,
      - . phenanthroline: end concentration 1 mM: 4 % relative activity.
- 5. Protein according to anyone of claims 1 to 4, characterized in that it is obtained in a purified state, from a cell extract of said *Lactobacillus delbrückii* ssp. *lactis* WS87 by the following procedure:
  - fractionation of the cell extract by salting out at 4°C with streptomycin sulfate,
  - centrifugation,
  - anion exchange chromatography by applying the supernatant obtained at the previous step to an appropriate column, such as a column of Q-Sepharose Fast Flow (Pharmacia), and pooling the eluted fractions having the highest specific activity against L-Pro-p-nitroanilide,
  - preparative isoelectric focusing, by applying the pooled fractions obtained at the previous step to an
    appropriate column, such as a LKB column (Types 8100-1), with ampholytes within a range of pH 4
    to 6, and pooling the eluted fractions having the highest specific activity against Lys-paranitroanilide.
- 35 6. Nucleic acid characterized in that it codes for a protein according to anyone of claims 1 to 5.
  - Nucleic acid according to claim 6, characterized in that:
    - it comprises all or part of the nucleic acid represented on figure 3, and by SEQ ID NO 1, coding for a protein according to anyone of claims 1 to 5, or its complementary sequence,
    - or it hybridizes with all or part of said nucleic acid represented by SEQ ID NO 1, or with its complementary sequence.
  - 8. Nucleic acid according to claim 6 or claim 7, characterized in that it codes for the PepN protein represented by SEQ ID NO 2, or for polypeptides derived thereof having said hydrolysing β-naphtylamides activity, said nucleic acid comprising all or part of the nucleotide sequence delimited by the nucleotide located in position 316 and the nucleotide located in position 2844 in the nucleotide sequence represented by SEQ ID NO 1.
  - Recombinant nucleic acid containing at least one nucleic acid according to anyone of claims 6 to 8, inserted into a heterologous nucleic acid.
  - 10. Recombinant vector comprising a vector sequence, notably of the type plasmid (such as plasmids originating from lactic acid bacteria), cosmid or phage, and a nucleic acid according to anyone of claims 6 to 8, in one of the non-essential sites for its replication, and optionally one or several nucleic acid(s) coding for other aminopeptidases such as PepX.
  - 11. Cellular host which is transformed by a recombinant vector according to claim 10, and comprising the regulation elements enabling the expression of the nucleotide sequence coding for a protein according to

anyone of claims 1 to 5, and optionally, other aminopeptidases such as PepX, in this host.

- 12. Cellular host according to claim 12, chosen from among bacteria such as *E.coli*, or lactic acid bacteria including *lactococcus* and, especially, lactobacilli of the thermophilic group, transformed by a vector according to the invention.
- 13. Process for preparing a protein according to anyone of claims 1 to 5 comprising the following steps:
  - the culture in an appropriate medium of a cellular host according to claim 11 or 12,
  - the recovery of the polypeptide produced by the above said cellular host from the above said culture medium.
  - the purification of the protein thus obtained.

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- 14. Fermentation process characterized in that it comprises a step of treatment of material to be fermented with:
  - an appropriate amount of the PepN protein or polypeptides derived thereof according to claims 1 to 5, and optionally other proteases, and more particularly aminopeptidases such as PepX, and/or
  - an appropriate amount of at least one of the transformed cellular host, according to claim 11 or 12,
  - and optionally, an appropriate amount of lactobacilli, such as Lactobacillus delbrückii ssp. lactis.
- 15. Process for the preparation of fermented foodstuff, and more particularly of cheese, which comprises a step of treatment of food material to be fermented, such as milk, with:
  - an appropriate amount of at least one of the protein according to claims 1 to 5, and optionally other aminopeptidases such as PepX, and/or
  - an appropriate amount of at least one of the transformed cellular host according to of claim 11 or 12, and/or
  - an appropriate amount of lactobacilli, such as Lactobacillus delbrūckii ssp. lactis.
  - 16. Process according to claim 15, characterized in that it comprises a step of treatment of food material to be fermented, such as milk, with other species and strains susceptible to be used as starter organisms in fermentation processes, and more particularly lactic acid bacteria susceptible to produce a PepX protein, such as Lactococcus, Streptococcus and Lactobacillus.
  - 17. Process according to claim 15 or claim 16, characterized in that it can be used as fermentation processes for the obtention of hard cheeses, such as Emmentaler type cheese
- 18. Foodstuff, and more particularly cheeses, such as obtained by fermentation processes, according to anyone of claims 15 to 17.

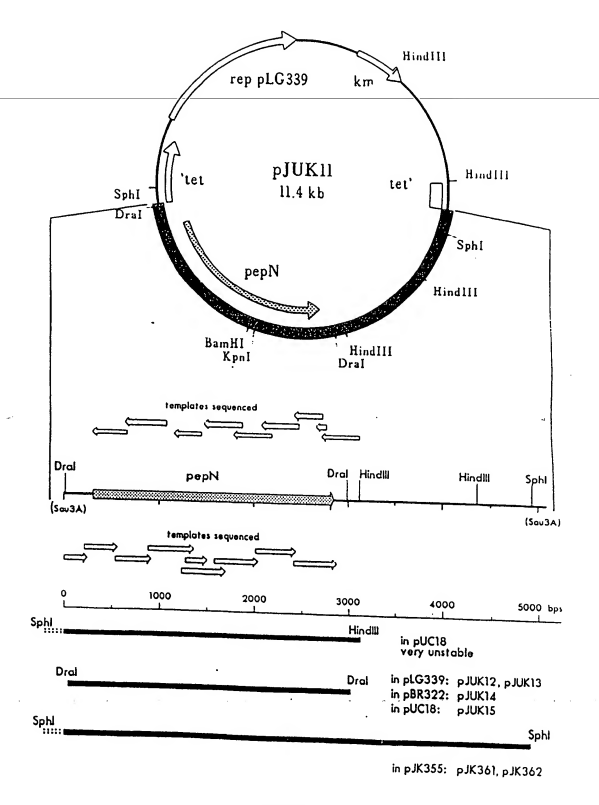


Figure 1

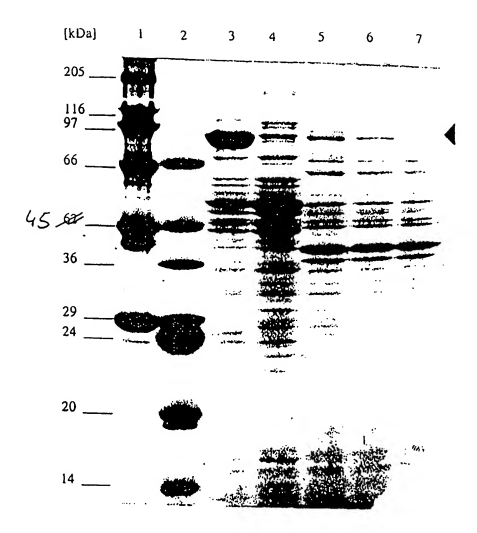


Figure 2

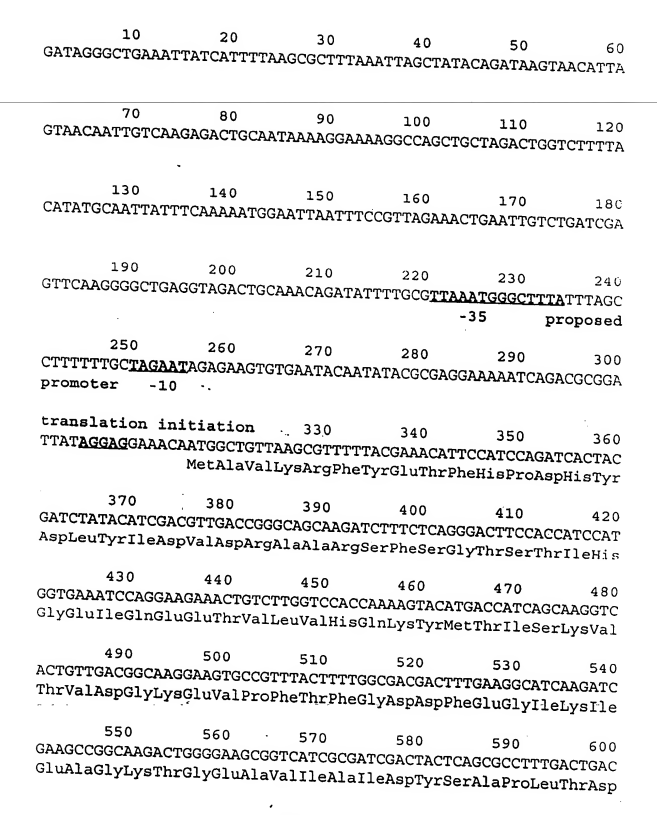


Figure 3

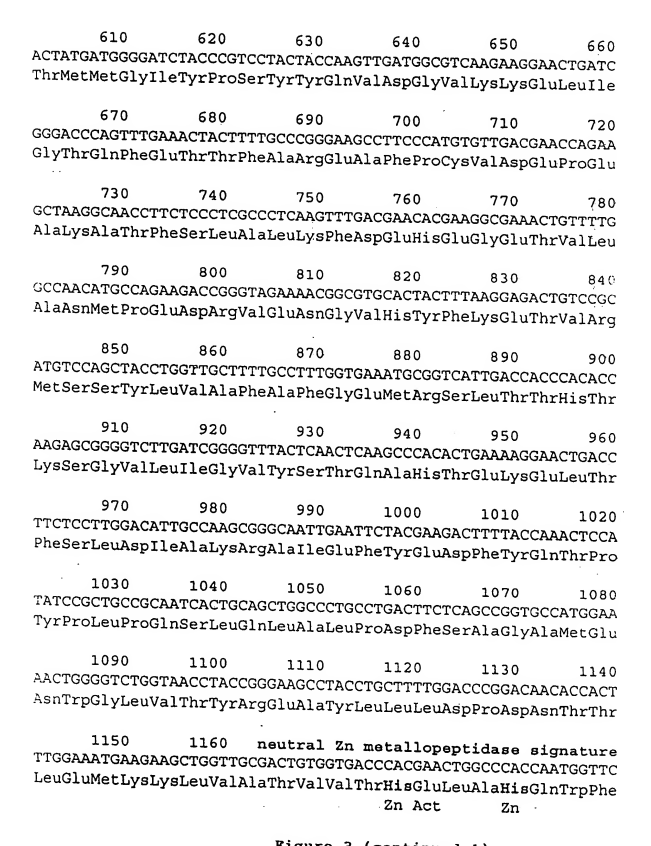


Figure 3 (continued 1)

1210	1220	1230	3040		
	CCATGGAATG	1230 CTCCC2C22	1240	1250	1260
GGTGACCTGGTAA	hrMotClm.	- Maria	CTCTGGCTGA	ACGAAAGTTI	CGCCAAC
GlyAspLeuValT	minecoluli.	prrpaspasn	LeuTrpLeuA	${ t snGluSerPh}$	eAlaAsn
1270	1280	1290	1300	1310	1320
ATGATGGAATACC	TGTCAGTTGA	CCACCTGGAA	.CCTAACTGGC	3.03.000000	
MetMetGluTyrI	euSerValAs	pHisLeuGlu	ProAsnTrpH	isIleTrnGl	11Met Dhe
			2		unecrie
1330	1340	1350	1360	1370	1200
CAGACTTCTGAAG GlnThrSerGluA	CAGCGGCTGC	CTTGACCCGG	באר א אינים שלים		1380
GlnThrSerGluA	laAlaAlaAl	aLeuThrArg	Acay Jampes	ACGGGGTACA	GTCAGTG
		garminig	reparatura	sbGTAATG1	nSerVal
1390	1400	1410	7400		
CACGTGGAAGTTA	ATGACCCGC	ייים אינים אינ אינים אינים אי	1420	1430	1440
CACGTGGAAGTTA HisValGluVala	engenbrog!	COMMATCGAC	GCCCTCTTTG.	ACGGGGCCAT	'CGTTTAC
HisValGluValA	GIWSDELOWI	aGIUITEAsp	AlaLeuPheA	spGlyAlaIl	eValTyr
1450	1460				
	146U	1470	1480	1490	1500
GCCAAGGGGTCAA AlaLysGlySera	GAATGCTGGT	CATGGTCCGG	TCACTTTTGG	GCGATGAAGC	CTTGAGA
AlaLysGlySerA	rgMetLeuVa	lMetValArg	SerLeuLeuG	lyAspGluAl	aleuara
				1 -1	ancantg
1510	1520	· 1530	1540	1550	1560
AAGGGCTTGAAGC LysGlyLeuLysA	GCTACTTTGA	CAAGCACAAG	שיי אי אי איי		1560
LysGlyLeuLysA	rgTyrPheAs	pLvsHisLvs	PheGlyAca	CGGCAGGTGA	CGATCTC
	_ <u>-</u>	1-1-11-11-11	THEOTY ASILA	raAraGryAs	pAspLeu
1570	1580	1590	7.500		
TGGGATGCCTTGT	CAACGGCCAC	TOPU	1600	1610	1620
TGGGATGCCTTGT TrpAspAlaLeuS	erThralamh	TGACTTGAAC	ATTGGGGAAA	TCATGCACAC	TTGGCTG
TrpAspAlaLeuS	CTIMIAIA	raspleuAsn	IleGlyGluI	${ t leMetHisTh}$	rTrpLeu
1630					
	1640	1650	1660	1670	1680
GACCAGCCAGGCT AspGlnProGlvT	ACCCAGTGGT	GAATGCTTTT	GTTGAGGACG	GCCACTTGAA	GCTGACT
AspGlnProGlyT	yrprovalva	lAsnAlaPhe	ValGluAspG	lyHisLeuLv	SLeuThr
					o D C Q I I I I
1690	1700	1710	1720	1730	1740
CAGAAGCAATTCT GlnLysGlnPheP	TCATCGGTGA	AGGCAAGGAA	ОШООООООО		1/40
GlnLysGlnPheP	helleGlyGl	uGlvLvsGlu	ValGlyarati	HOTOGOMAAT	TCCGCTT
	-	2-10010	- ~ TOT YAT GL	AsitbGInII	eProLeu
1750	1760	1770	1700		
AACGCTAACTTCA AsnAlaAsnPhet	AGGCACCGAA	₽₹₹₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽	1780	1790	1800
AsnAlaAsnPheL	vsAlaDrot.	off cwickled (A)	SACGTTGAAC	TTGACCTGGG	TGACTAC
	1 - WITGELOUD	erremerger!	AspValGluLe	Countrakus	VA en Tur

# Figure 3 (continued 2)

. — •	1820		1840	1850	1860
CAGGCTCTGCGGG	CAGAAGCCGG	CCACGCTCTG	CGCTTGAACG	TGGGCDACNN	CTCCCAC
GlnAlaLeuArgA	laGluAlaGl	yHisAlaLeu	ArqLeuAsnV	alGlvAsnas	ncarlia
			J= J=	arerynsims	MOCINIS
1870	1880		1900	1910	1920
TTCATCGTGAAGT	'ACGACCAGAC	TTTGATGGAC	GACATCATGA	ACCA ACCCA A	CC r commo
PhelleValLysT	yrAspGlnTh	rLeuMetAsp	AspIleMetL	ysGluAlaLv	sAspLeu
1930	1940	1950	1960	1970	1980
GATCCAGTTTCCC	AATTGCAATT	GCTGCAAGAC	CTGCGGCTTT	TGGCAGAAGG	CARCONA
AspProValSerG	SlnLeuGlnLe	uLeuGlnAsp	LeuArgLeuL	euAlaGluGl	yLysGln
1990	2000	2010	2020	2030	2040
GCTTCATACGCTG	ACGIGGIACC	AGTTCTGGAA	CTCTTCAAGA	ACTCAGAAAG	CCACATT
AlaSerTyrAlaA	spvalvalpr	oValLeuGlu	LeuPheLysA	snSerGluSe	rHisIle
2050					
	2060	2070	2080	2090	2100
GTCAACGATGCTC	.IGTACACGAC	TGCTGATAAG	CTGCGGCAAT	TTGCCCCAGC	CGGCAGT
ValAsnAspAlaL	euryrnnri'n	rAlaAspLys	LeuArgGlnP	heAlaProAl	aGlySer
2110		•			
	2120	2130	2140	2150	2160
GAAGCTGACAAGA GluAlaAspiyaA	Splany~~yl	TCTGTACAAC	GACTTGTCCA	AGGACCAAGT	TGCCCGT
GluAlaAspLysA	ratine dat dat	aLeulyrAsn	AspLeuSerL	ysAspGlnVa	lAlaArg
2170	2180	21.00			
· ·		2190	2200	2210	2220
TTGGGCTGGCTGC LeuGlyTrpLeuP	rolveyladi	vejuse-ye.	GAAGACATTC	AGACCCGGCC	ATACGTT
1	LODIBRIAGI	yGluSerAsp	GluAspileG	lnThrArgPr	oTyrVal
2230	2240	2250	2260		
TTGTCTGCTAGCC		CAACGCTGaT	226U TCDGDDDDCC	2270	2280
LeuSerAlaSerL	euTyrGlyAr	GASDAlaAsD	serglulvec	AAGCCCACGA	AATCTAC
	•	J	ocidianysu	THATAHISGI	ulleTyr
2290	2300	2310	2320	2330	2242
GTGGAATACGCTG	ATAAGTTGGC	AGAACTGTCC	GCTGATATCC	<b>ににていかれることか</b>	2340
ValGluTyrAlaA	spLysLeuAl	aGluLeuSer	AlaAspTleA	COCCATACGI CODCOTURUS	lingale
				- 3. LOIYI Va	THEUITE
2350	2360	2370	2380	2390	2400
AACGAAGTTGAAA	ACTACGGGTC	AAGCGAATTG	ACTGACAAGC	ጥር እ ጥጥር <u>ር</u> ጥጥጠ	CMACOAC
AsnGluValGluA	snTyrGlySe	rSerGluLeu'	ThrAsplivet	enllector.	OTACCAG
	-				arArgru

Figure 3 (continued 3)

2410	. 2420	0420	_		
	. בעבט ירשתית אחת שרי	2430	2440	2450	2460
GCAACCAGTGACC	CAICAIICA CAICAIICAI	AGATGGACCTG	GAAGCCGCGA	TCGTGAAGAG	CAAGGAC
AlaThrSerAspP	roserbueri	ysmetAspLeu	GluAlaAlaI	leValLysSe	rLysAsp
2470	2480				
	2480	2490	2500	2510	2520
GAAGGCGAACTGA GluGlyGluLeuI	AGAAGATCG:	LTTCCTGGTTC	AAAAACGCTG	AAATCGTTAA	GCCGCAG
GluGlyGluLeuL	AarAatteA	alSerTrpPhe	LysAsnAlaG	luIleValLy	sProGln
2530				_	
	2540	2550	2560	2570	2580
GACTTGCGCGGCT AspLeuAraGlyT	GGTTCAGCG(	SCGTTTTGTCC	AACCCGGCAG	GTGAACAGCT	GGCCTGG
AspLeuArgGlyT	rpPneSerG.	lyValLeuSer	AsnProAlaG	lyGluGlnLe	uAlaTro
2590					
	2600	2610	2620	2630	2640
GACTGGATCAGAG AspTrplleArgA	ACGAATGGG(	CATGGTTGGAA	AAGACGGTCG	GCGGCGACAT	
AspTrpIleArgA	spGluTrpA]	laTrpLeuGlu	LysThrValG	lyGlvAspMe	tGluPhe
				1 1 1	COLUPILE
2650	2660	. 2670	2680	2690	2700
GCTACCTTCATCA AlaThrPheIleT	CTGTCATCT	CCCGCGTCTTC	AAGACCAAGG		2700 CCAATAC
AlaThrPheIleT	hrValIleSe	rArgValPhe	LysThrLysG	luaraturas	DATABOO
			1	runigiyins	pGIuTyr
2710	2720	2730	2740	2750	07.45
AACGCCTTCTTTA AsnAlaPhePheT	CTGACAAGGA	AAGCAACATG	00000000		2760
AsnAlaPhePheT	hrAspLysG]	uSerAsnMer	Lentengery	GGGAAATCAA ~~ClXlX	GATGGAC
	_		CCADCARSIM	ragratterA	sMetAsp
2770	2780	2790	2800	2070	
CGGAAGGTCATCG ArgLysVallleA	CTAACCGGGT	ישיתי עבותיים עבועי	70020022	2810	2820
ArgLysValIleA	laAsnArgVa	llAspLeulle	Alasarciuc	AAGCTGACGT	CAACGCC
			er aber Grag	Inalaaspva	lAsnAla
2830	2840	2850	220-0		
GCGGTTGCTGCTG AlaValAlaAlaA	CTTTGCAAAA	GTAATTCAAT	odorų Davadonos	sed transc	ription
AlaValAlaAlaA	laLeuGlnLv	SEnd	AGAGCATAAG,	AAAACTG <u>TT</u> T	CCGCTGA
	7	<u>-</u>			
2890 ·	2900	2910	2020		
GAGCTGGAAACAGGGGAGGGGAGGGAGGGGAGGAAACAGGGAAACAGGGAAACAGGGAAACAGGAGG	TTTTTTTATG	ምምም ለ ለግሞው ያቸው። የጥውጥ ለ የግሞው ያቸው	2920	2930	2940
terminator			FICIGCAATC(	GGTTACAATA	TAGATGT
2950	2960	2070	0055		
	┸₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽	2970	2980	2990	3000
AAATACTATCGTA		GOTAATAAAA	'GAACAACGA'	TTTTAAAGAT	ATCATGC

Figure 3 (continued 4)

3010 3020 3030 3040 3050 3060 AAAACAGAAAGTCTATCCGGCACTATGATTCCAGCGTGAAGATTTCCCGTGACGAATTGC

3070 3080 3090 3100 3110 3120 TGGAAATCATTAATGAATCTATCTCTGCTCCAAGTGCCTGCAACCTGCAGTCCTGGAAGC

TT

Figure 3 (continued 5)

	OKGAN I SM	REFERENCE	MOMOLOGY in complete protein sequence Ab (conserved amino acid changes permitted)	Abreviation in alignment	ent
LLLYSAP CICKHINO CICKHINO CICKHINO HSANDEPH MASATIO MASATIO ECPEPH ECPEPH	Lc. factis ssp. lactis MG1363 Lc. factis ssp. cremoris Wg2 Trabbit Momo sapiens rat mouse Saccharomyces cerevisiae	Tan et. al. 1992a Stréman 1992 Noren et al. 1989 Helmiss 1989 Uu et al.1990 Garcia-Alvarez et al. 1991 McCanan and Gabe 1986	62 x 62 x 1c. 1c. 41 x 1st	Lc. lactis Lc. cremor rabbit man rat mouse S. cerevis	
Lc. lactis Lc. cremor Lb. lactis S. cerevis S. cerevis ran rat rabbit mouse E. coli	My FAEEE	WAAVCTIIALSVYYSGEKIKRANSSPYAST WAAVCTIIALSVYYAGEKIRRAENSATAPT VGLIVGLSVGLTRSCEGOTTPAPSGPPPEA	MAVKR LIETFVPENYKIF LDIDRKTKK IKGGVAIT MAVKR LIETFVPENYKIF LDIDRKTKK IKGGVAIT MAVKR LIETFVPENYTIF LDIDRKTKK IKGGVAIT MAVKR FYETFHPDHTOLT IDVORAARS FSGTSTIF MAKGFYI SKILGIL GILLGVAAVCTIIALSVYTSOEKHKHANSSPVASTTPSASAŤÍNPASATTLDOSKANRYRLPHILKPÖSTGVT LRPYLTPHORGLYVFKGSSTVR MAKGFYI SKTLGIL GILLGVAAVCTIIALSVYTAGEKHRHAEHSATAPTLPGSTSATTSTTHPAIDESKPUNOVRLEVILIPDSTGVT LRPYLTPHEGGLYTFKGSSTVR MHFAEEEPSKKYCIKGKHVAIICGVVVAVGLIVGLSVGLTRSCEGOTTPAPSOPPEASTALPPGDGHVCPDSEDESGEKKHRLPDFINPVHYDLE VKALMEEDR YTGIVTIS	IKGOVAIT IKGOVAIT FRGTVAIT FRGTVAIH FEGSWIE ALVFKGSSTVR ALTIFKGSSTVR YTGIVTIS	
Lc. lactiss Lc. cremor Lb. lactis S. cerevis man rat rabbit mouse E. coli Lc. cremor Lb. lactis S. cerevis man	GGAKO TVVSFHTKGLI GGEAKO TVAFHKAKGLI GGEAKO TVAFHKAKGLI GGEAKO TVAFHKAKGLI LKI NNPAIDTYTLATVOT FTCKETT NVI I I INSKKLIN FTCKETT NVI I I INSKKLIN FTCKETT NVI I I INSKKLIN WALSKPT ROLULHI RETK GGEOFAKAT FOLSVK FOE SIDEFEKAT FOLSVK FOE SIDEFEKAT FOLSVK FOE CVOEPEKAT FSLALK FOE CFDEPAKAS FNI I I. I HPN CFDEPAKAS FNI I I HPN CFDEPAKAS FNI I I. I HPN CFDEPAKAS FNI I HPN CFDEPAKAS FNI I HPN CFDEPAKAS FNI I T HPN CFDEPAKAS	GEAKO TVVSFHTKGL HFUKVRAFSVOTNFIENEEDEETV GEAKO GEAKO TVASHAKGL HFUKVRAFSVOTNFIENEEDEETV GEAKO TVASHAKGL HFUKVRAFSVOTNFIENEEDEETV IEGETGE ETVLVHQYTHTISKYTHOOKGEVPFTFADOFEGIV IEGETGE ETVLVHQYTHTISKYTHOOKGEVPFTFADOFEGIV IEGETGEVT FAFFKGHAFF FTCKENT DVIIIHSKYLNYTLSGGRRVALRGGGSOPPDIDKTELVEPTEYLVMHKGSLVK GHOYE FTCKENT DVIIIHSKYLNYTLSGGRRVALRGGGSOPPDIDKTELVEPTEYLVMHKGSLVK GHOYE WALSKPT ROLULHIRETKITKLPE LRRPSGGOPDIDKTELVEPTEVLOSSLVK GHOYE WALSKPT ROLULHIRETKITKLPE LRRPSGGOPDIDKTELVEPTEVLOSSLVK GHOYE WALSKPT ROLULHIRETKITKLPE LRRPSGGOPDIDKTELVEPTEVLOSSLVK GHOYE WALSKSTAITLIHPE LRRPSGGOPWAKKEFNITLIHPK TTALSHALPKOPSTPLPEDPHANYTEFTTRONSSTLAFVGGLOYKK FOEPAKKASFNIILIHPK FOLTALSHALPKOPSTPLPEDPHANYTEFTTRONSSTLLAFVGGLOYKK FOEPAKKASFNIILIHPK FOLTALSHALPKOPSTPLPEDPHANYTEFTTRONSSTLLAFVGGLOYKK FOEPAKKASFNIILIHPK FOLTALSHALPKOPSTPLPEDPHANYTEFTTRONSSTLLAFVGGLOYKK FOEPAKKASFNIILIHPK FOTALSHALPKOPSTPLENGHY FOLTALSHALPKOPSTPLPEDPHANYTEFTRONSSTLLAFVGGLOYKK FOEPAKKASFNIILIHPK FOTALSHALPKOPSTALHTPRONSTLLAFVGELOYKE FOEPAKKASFNIILIHPK FOTALSHALPKOPSTALHTPRONSTLLAFVGELOYK FOEPAKKASFNIILIHPK FOTALSHALPKOPSTALHTPRONSTLLAFVGELOYK FOTALSHALPKOSTALHTPRONSTLLAFVORGOLVH FOLDALPFAGAREHWGLYTYREVCHLUDDPHANTYTEHKLANOFFGLLYHE FOLDALPFAGARHWGLYTYREVCHLUDDPHANTYTEHKLANOFFGLLYHE FOLDALPFAGAARHWGLYTYREVCHLUDDPHANTYTHELAHOWFGLLYTHEMDHLUNE FOLDALPFAGAARHWGLYTYREVCHLUDDPHANTYTHELAHOWFGLLYTHEMDHLUNE FOLDALPFAGAARHWGLYTYREVCHLUDDPHANTOFFHLHFELAHOWFGLLYTHEMDHLUNE FOLDALPFAGAARHWGLYTYREVCHLUDDPHANTYTHELAHOWFGLLYTHEMDHLUNE FOLDALPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRESALVENDHUNG FOUGLADPFAGAARHWGLYTYRENDHUNG FOUGLADPFAGAARHWGLYTYRENDHUNG FOUGLADPFAGAARHWGLYTYRENDHUNG FOUGLADPFAGAARHWGLYTYRENDHUNG FOUGLADPFAGAARHWGLYTYRENDHUNG FOUGLADPFAGAARHWGLYTYRENDHUNG FOUGLADPFAGAARHWGLYTYRENDHUNG FOUGLADPFAGAARHWGLYTYRENDHU	GEAMD TWYSFITKGLWENKYRAFSYDTWFIEWEDEELY WIGETORYT GEAMD TWYSFITKGLWENKYRAFSYDTWFIEWEDEELY WIGETORYT GEAMD TWYSFITKGLWENKYRAFSYDTWFIEWEDEELY WIGETORYT GEAM TWYSFITKGLWENKASHOTWFIEWEDEELY WIGETORYT GEAM TWYSFITKGLWENKASHOTWFIEWEDEELY TEACH TO THE STANDARD THE STANDAR	GFESHFARGAFP  GFESHFARGAFP  GFETTFAREAFP  GHAADARKSFP  GHGAADARKSFP  GHCAADARKSFP  TEDTTOTPTPIP  TEDTTOTPTP  TEDTTOTPT  TEDTTOTPTP  TEDTTOTPT  TEDTTOTPT  TEDTTOTPT  TEDTTOTPT  TEDTTO	
794	KLOKIA POGETAMENGUVI YRETHLYDELLSASSHOORVASVUTAMELANGNEGNU TVOMMNDLLLNEGFASVE IYM IVAVOFFNHGAMENGLVI YRETHLI YDPLLSASSHOORVASVVAHEL VHOWFGNI VI MUMADDLULNEGFAST FEI IXM IVAVOFFNHGAMENKGLNI FNSKYVLARTDTATOKOVLOFERVIGHEY FNNHTGNRVTCROUFOLSLKEGLI VFRO	PTLYSSISMEKYVI I VAMELAHOWFGHLV PPLLSASSHOORVASVVAHEL VHOWFGHTV: TDTATDKDYLDIERVIGHEYFHWYGMRVI	KLDKIATOP GEGAERHUGL VYYRETHLI YDPLLSASSHOGNYGAN VAMFLAMGAFGKIVY VOLKHOL HLEGFASYVEYL GADYAEPTWILKDLI VLNEI HSVMAVDALASSHPLSSPA IYM IVÄVDFFKHGAMEHKGLUT YRETHLI YDPLLSASSHOGNYGYVAHELYHOUFGNTY HDUMDDI ULNEGFAST FEFI GVNHAEKDUGHLSGVLUSDEDSLHSSHPV IYM IVÄVDFFKHGAMEHKGLNI FNSKYVLARTDTATDKDYLD IERVI GHEYFHHUTGARVI CROUFGLSLKEGLI VFRQ GEFSSLGSRAVNRI NHVRTHRGLGFAEDASPM AHP	SSHPLSSPA SSHPV V DASPM AHP	

Figure 4

VAAFHHSUVNOPGPPUVT AEVVDDILLISGKGFFVGEGVDK VAAFHHSUVNOPGPPUVT AEVVDDILLISGKGFFVGEGVDK VAAFHHSUVNOPGPPUVT AEVVDDILLISGKGFFVGEGVDK VRSVHNIJVTKKVGFPVISVEBGHGKLTFRQHFLLDPDSNV VSFI MORHILQHGFPVIT VDTSTGTLSGHFLLDPDSNV VKEVMDILLQHGFPVIT VNTSTGEIYGEHFLLDPTSKP VKD IMDRILLQHGFPVIT VNTSTGEIYGEHFLLDPTSKP VKEVMDILLSGGFPVYT VYSGEI NITQKFLLDFTSNV VKEVMDILTSGHGFPVYT VYGDYNPETEQYTLTISGRT	HWAKY I DYKGALL TOLLSEVESLENYTKFOILODRKLLAKAGVISYADVWILPSFTNEE NYAHT I DYKGALL TOLLSEVESLENYTKFOILODRKLLAKAGVISYADVWILPSFTNEE NISHFI WYTDOTLH DO INKEAKOLDPVSQLOLOD RILAEGKQASYADVPVIELFKNSE SSGIY I TSYTDERW AKLOGADLLSVEDRYOLVADVKILSASGYTSTNFLNLVSKANNEK VTGYTRVNYDEENWRKIGTQLORDHSAIPVINRAGI INDAFNLASARKPYTLALNINTFLIEER YTGYTAVNYDEGNIKKIGTQLOTDLSVIPVINRAGI INDAFNLASAGKLSITLPLSNTLFLASET YTGYYQVNYDEGNIKKLGTQLGTNPSVIPVINRAGI INDAFNLASAGKVPYTLALDNTLFLIRET HIGFYRWYTEGGTLÖNIAEALSSNHTRFSAADRSSFIDDAFALARAGILIKKIALLINTKESE WYTFGP WYTFGP WYTFGGTLÖNIAEALSSNHTRFSAADRSSFIDDAFALARAGILIKKIALLINTKESE	SOIFAAH KE NLASIPAD IRPIYLMNEIKTIUS SDIFATH KE NLASIPAD IRPIYLMNEIKTIUS KETYVET AD KLAELSAD IRPIYLINEVERYGS KETYVET AD KEALSAD IRPIYKIINEVERYGS GLYGOWHONPIHPH LRSTYYCHAIA GG GLYSOWHNSONNPIHPH LRSTYYCHAIA SC DLFKOWDDPSHNPIHPH LRSTYYCHAIA GG GLFSOWHUSONNPIHPH LRSTYYCHAIAL GG GLFSOWHUSONNPIHPH	LEEKLGGDMS YOKFVIYPGNIFKTAD KLA LEKLGGDMS YDKFVIYPGNIFKTAD KLA LEKYVGGDME FAFFIYISRVFKTKE RYD LVXRLPPGLSWLGSVTLGTSGFTSMO KID PFDYTGGGSFSFSWLJOGYTRRFSTEY ELG LFEDYGGGSFSFAWLJOGYTRRFSSEF ELG LFEDYGGGSFSFAWLAGVTRRFSSEF ELG LFEDFGGGSFSFAWLIAGVTRRFSTEY LVSRFTIMD RYLGRIVTIAEPFHTEL QLU ISLIGAFAGSWPAAFHAEDGSGYLFLVEWLTDLN	
VEVTHPDE I GIL FDPA I VTAKGSRL MYHLRKUL GDED FAAGLALT FRRHGTGHI VGDHLUDALAEVSGKD VAAFHHSLVNAOPPEVT VEVUPPE I GIL FDPA I VTAKGSRL WYHLRKUL GDED FAAGLALT FRRHGTGHI VGDHLUDALAEVSGKD VAAFHHSLVNAOPPEVT VEVUPPEL IDAL FDGA I VTAKGSRL WYHLRKUL GDEALRGL KRYFDKHKFGNAAGDOL MDALSTATDLH VEVKKADE I NOI FD AI SYSKGASLL RHITSKLIGEET FIKGVSOYL HKFKYGNAKTEOL WALLADASGKE VRYTKADI SEL FD AI SYSKGASLL RHITSKLIGET FIKGVSOYL HKFKYGNAKTEOL WALLADASGKE VRYTKADI SEL FD AI SYSKGASLL RHITSKLIGE FIT FOR SYNTI TILLUMHLOANNSOYAI KLPASYST HORHIL GHGFPVIT DEWNIPPAISEL FD SITTSKGASVL RHILSFLIFEDL FKGGLASYLHT FAYONTI YLDLWEHLQAANSOSAI QLPASYNDI HORHILGHGFPVIT VLYSTPAEITSVFD GISYSKGASILRHL GOUITTEKFRGKGOTI YLKFRFORMATSOFWANKEDASNV LSHFRRHYSGSGFPIV LSHFRRHYSGSGFPIV	LUNV PLNINTGLPOL LSSEKVEIPGFAALKTKNHGKALFLNDANHAHYIIDYKGALL LWAV PLNINTGLPOL LSSEKKEIPGFAALKTKNHGKALFLNDANHAHYIIDYKGALL KWEI PLNINFKA PKI HSDVELDLGDYGALRA EAGHARILNYGANSHFUKTGOLL DKTITPVFLALKTKNGV DSSVYLSERSKTIELEPPTFFKVNSEGSGIYIISYTDERU VNIV PITSIRDGRQQODYLLUD VRAGNDLFSTSG NEWLLLHINVTGYTONNYDEGNURSIPVINRAGIINDAFNLASAKTVPTLALHNITLELIEST LWIV PIPYLKNGKEDH YULET EKKOSAEFOTSS NEWLLLHINVTGYTONNYDEGNURKIGTQLGNDHSAIPVINRAGIINDAFNLASAKTVPTLALHNITLELIEST LWIV PYSSKHSVGOOGFFLLEGVEGTONSLFRVEGDNHWILLAMINTGYTONNYTGENKKLOTOLGNDFSTPVINRAGIINDAFNLASAGKLSITLPLSNTLFLIRFT TANI PVRAKADNDANSRITYY NRLDKGGITLMANLSGDFLKINPOHIGFYVPVEGTKONIAAGALSSHHTRFSAADRSSTIDDAFALARAGILHYKLSEE QPLHIPFAIELYDNEGKVIPLQKGGNPVNSVLWYTGAEGTFVFD VVALLEFSAPVKLEYKNSDOOLTFLNRHARNDFSRUDAAGSLLATYKLNVRHO	SYLVNTGLSQLISELE LFVDEDSETEKAFOSLVGKLFAKNYARI SYLVNTGLSQLISELE LFVDEDSETEKAFOSLVGKLFAKNYARI GADVAGESAGDESLRGIVLSKTLYSENADAKTKASOIFAAH KE NLASIPAD IRPIVLNNEIKTINS SHIVNDALYTTADKLR GAPAGSEADKNLEALYDLSKDOQVARI SHIVNDALYTTADKLR GAPAGSEADKNLEALYDLSKDOQVARI GALPKAGESDEDIOTPYVLSSLYGRANDSETOKAHETYYK AD KLAELSAD IRPIVLNYRVBKNGGS SFVYADOIINSISSHKSTUFFEKETOALDNFTKKLISGATHHL GALPKAGESDEDIOTPYVLSSLYGRANDSETOKAHETYYCH AD KKAIPAL IRPIVLNYRVBKNGGA STYVADOIINSISSHKSTUFFEKETOALDNFTKKLYTHRGARGARGAARDADVEKAALKYTOKGARDHINPIHPH LRSTVYCHAIAS GE SYNDAMALSSLYTISHE DRSEVYGPHKNYLSKOYPLFEHFKNITMANTRIP DILMCOYNEINASTGSGLEECROLVGLYSOMHNSONNPIHPH LRSTVYCHAIAS GE SYNDAMALSSLYTISHE DRSEVYGPHKNYLSKOYPLFEHFKNITMANTRIP DILMCOYNEINASTGSGLEECROLVGLYSOMHNSONNPIHPH LRSTVYCHAIAI GG GA QD GSHITKLIRASIIGFECKCHONGOPSHROIHPH LRTVYCHAIAL GG GGOPLSIPVHVADAFRAVLLDEKIDDEKINDALAGEITLIPSVHEMAELF DIIDPIAIAEVREALISTLATELADELLAIYNANYGSETRVEHEDIAKRILRHAGLRFLAF GETHLADV	AELVKTYRETYIKTSLOEFKRELEGAVALIKDEKVIAELLESFKWADIVKPO  SELINKTYRETYIKTSLOEFKRELEGAVALIKDEKVIAELLESFKWADIVKPO  SELINKTRETYIKTSLOEFKRELEGAVALIKDEKVIAELLESFKWADIVKPO  SELIDKIIGLYOATSOPSFKADLEANIKSKOEGELKKIVSHFKWAEIVKPO  ENYEKYYK IYLDPISNDEKLAALRSLGRFKEPKLLERTICYLFOGYVIN  EEEMDFAMEOFRAATIVNEADKLRSALGSREUIHRYLSYTLHPOLIRKO  EEEMDFAMEOFRAATIVAEALKOEGSFSFSHIJOATRRFSTEY  DATST IVSIANNYVGOTLVADFVRSHWK  LEDYGGGSFSFSHIJOATRRFSTEY  LYSKQFHEANNHTLEOFOKTSLAGEKELLYGLARYLERTLERTLORUINKTOPHIKTO  LYSKTIND RYLGRIYIAETEY  LYSKQFHEANNHTDALAALSAAVAAGLPCROALLARYLERTLORUINKTOPHIKTO  LYSKGFHEANNHTOLAGSTURPANTIECTERTURD RYTIAELTRANDER AND ASTAL INGERORUSPANTELETRANDER AND ASTAL INGERORUSPANTELETRANDER AND ASTAL INGERORUSPANTELETRANDER AND ASTAL INGEROBOSCYLEVEHLERTELETRANDER AND ASTAL INGEROPORTICELETRANDER AND ASTAL INGEROPORTICE AND ASTAL INGERTANDER AND ASTAL INGERNATION AND ASTAL INGER AND ASTAL INGER AND ASTAL INGER AND A	EYKAFFEPKLEN GGLKRSIEMAIKQITARVALIDSGKAAVDKAITDIAEKL EYKAFFEPKLEN GGLKRSIEMAIKQITARVALIDSGKAAVDKAITDIAEKL EYNAFFTDKESN MLLNRE KWORKVIANRVALISGADAVOKAITDIAEKL EYNAFFTDKESN MLLNRE KWORKVIANRVALIASGADAVARAAALOK EIKKFFATK ST KGFDGSLAGSLOTITSKAGVAVNRDRDVANKYLKENGTY GLEGFKDDNESTGFGSGTRALEGALEKTKANIKAVKENKEVALGNFTENS OLEGFKEDNSATGFGSGTRALEGALEKTKANIKAVKENKOVALKAFTEN GLEGFKEDNATGFGSGTRALEGALEGTRANIKAVGENKEAVLANFTANSA ONGSFFKYPNA GAGARCREGALEGTVKNNIEHLINVNROSIREWFASLP SRNPQVASRLIEPLIRLKRYDAKRGEKHRAALEGLKGLENLSGDLYEKITKALA
LC. lactis VEVTHPDEIGILFDPAIVYA LC. cremor VEVTHPDEIGILFDPAIVYA Lb. lactis VEVMDPAEIDALFDGAIVYA S. cerevis VPVKXADEINOIFD AISYS man SEINTPAGISELFD AISYS rat NEVNTPAGISELFD SITYS rabbit DEVNTPAGISELFD SITYS mouse VTVSTPAEITSVFD GISYSI E. coli IRPOHVIEMNNFYILIV YEI	Lc. lactis GR LLMN PLHTNUT Lc. cremor GR LLMN PLHTNUT Lb. lactis GR KWEI PLNANFR S. cerevis DE DKITPVFLALK man TR PSEMYVLIV PIPYLK rabbit TR PSDFWTLUIV PIPYLK mouse SQPPSELGTTNNI PVSKNKN mouse SQPPSELGTTNNI PVSKNKN E. coli PATPQQAEKQPLHIPFAIELY	LC. lactis SYLVNIGLSQLISELE LFV0I LC. cremor SYLVNIGLSQLISELE LFV0I Lb. lactis SHIVNDALYTTADKLE GFAPI S. cerevis SFVVADQIINSISSMKSTWLF man OYNOWEAALSSLSYFKLMF rat EYMPWEAALSSLNYFKLMF rabit EYMPWAALSSLNYFKLMF mouse DFLWERYISSNSYIISMF E. coli QGQPLSLPWYADAFRAVLLOE E. coli	LC. lactis AELVKTYRETYIKTSLGEFKE LC. cremor AELVKTYRETYIKTSLGEFKRE Lb. lactis SELTDKLIGLYQATSDPSFGDO S. cerevis ENYEKYYK IYLDPISNDEKLA man EERDFAHEGFRAATLVNEADKI rabbit EERANFAWATVPERTLVNEADKI rabbit EERANFAWATVPERTLVNEADKI mouse EAAMNTHEGYGKTSLAGEKEKE E. coli LVSKQFHEAMNTDALAALSAA	Lc. lactis EYKAFFEPKLEN GGLKRSIEN Lc. cremor EYKAFFEPKLEN GGLKRSIEN Lb. lactis EYNAFFTDKESN MLLUNEIKH S. cerevis EIKKFFATK ST KGFDGSLAG man GLEGFKEDNETGFGSGTRALEG, rabbit GLEGFKEDNSTGFGSGTRALEG rabbit OLGFRLNNLDTGFGSGTRALEG mouse OMGSFFAKPNA GAGAKPREN E. coli SRNPOVASRLIEPLIRLKRYDAKI
		Figure 4 (c	continued 1)	_ <b></b>

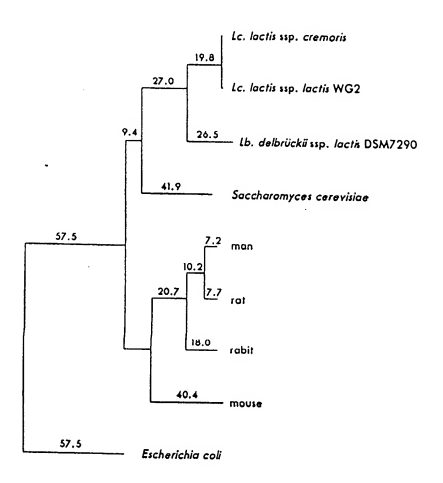


Figure 5

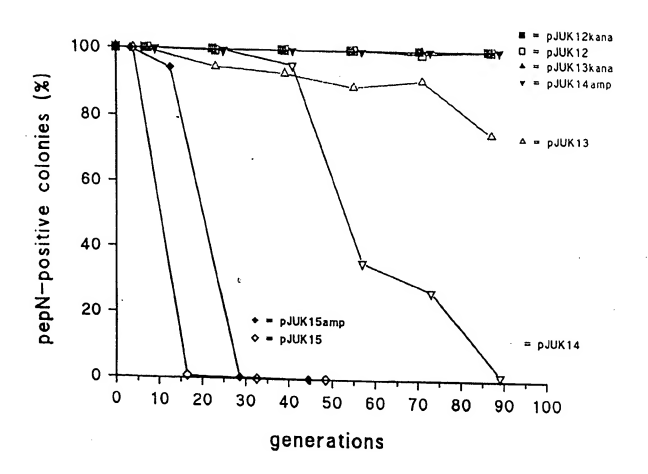


Figure 6



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]	DOCUMENTS CONSI	DERED TO BE RELEVAN	T	
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·	Place of search	Date of completion of the search	<del> </del>	Exception
	THE HAGUE	14 October 1994	. V	an der Schaal, C
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ategory	Citation of document with in of relevant par		Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CL6)
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Y	EP-A-0 522 203 (PRO January 1993 * abstract *	BICOM RESEARCH B.V.)	14-18	
<b>₽,</b> Y	CHEMICAL ABSTRACTS, 9 May 1994, Columbu abstract no. 237307 E. MEYER-BARTON ET sequence analysis of X-prolyl-dipeptidyl (pepX) from Lactobalactis DSM7290 page 251; abstract * & APPL. MICROBIOL. vol.40, no.1, Octob pages 82 - 89	s, Ohio, US; AL 'Cloning and of the -aminopeptidase gene cillus delbrueckii ss	1-18 5.	TECHNICAL FIELDS SEARCHED (Int. Cl.6)
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	of relevant passa	\$cs	to claim	APPLICATION (Int.CL6)		
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	* the whole document	*				
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-, ·	The present search report has been	drawn up for all claims				
	Place of search	Date of completion of the search		Examiner		
	THE HAGUE	14 October 199	4 Van	der Schaal, C		
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